

# SAMPLING AND ANALYSIS PLAN (FIELD SAMPLING PLAN AND QUALITY ASSURANCE PROJECT PLAN) JULY 2013

DATA GAPS ASSESSMENT TANK FARM 2, CATEGORY 1 AREAS NAVAL STATION NEWPORT PORTSMOUTH, RHODE ISLAND

PREPARED FOR:
NAVAL FACILITIES ENGINEERING COMMAND MID-ATLANTIC
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PREPARED UNDER: CONTRACT NUMBER N62470-08-D-1001 "CLEAN" CONTRACT TASK ORDER NO. WE30 Project-Specific Sampling and Analysis Plan Site Name: Tank Farm 2 Project Name: NAVSTA Newport Site Location: Newport, Rhode Island

Title: Data Gaps Assessment Document No.: W5211722D Revision Number: 0 Date: February 2011

# Worksheet #1 - Approval Page (UFP-QAPP Manual Section 2.1)

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Document Title:	Draft Sampling and Analys Project Plan), February 20 Newport, Newport, Rhode	is Plan, (Field Sampling Plan and Quality Assurance 11, Data Gaps Assessment, Tank Farm 2, Naval Station Island
Lead Organizat	tion: Naval Facilities Engin	eering Command Mid-Atlantic
Preparer's Nam	ne and Organizational Affili	ation: Tetra Tech NUS, Inc.
Preparer's Add	ress and Telephone Number	er: 234 Mall Boulevard Sulte 260, King of Prussia, Pennsylvania 19046-1433
Preparation Date	te (Day/Month/Year): Febru	ary 2011
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Investigative Org Manager:	anization's Project QA	It & Johnston 2-4-2011
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Lead Organizatio	n's Project Manager.	Cional and Date
		Signature/Date
		Roberto Pagtalunan, PE, NAVFAC Mid Atlantic
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Lead Organization	n QA Officer:	DN: c=US, o=U.S. Government, ou=DoD, ou=PKI, ou=USN, cn=80WERS.KENNETH.A.1230092474
		Date: 2011.02.23 13:31:52 -05'00'  Signature/Date
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		Signature/Date ·
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Signature/Date

Gary Jablonski, RIDEM

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Title: Data Gaps Assessment Document No.: W5211722F Revision Number: 0 Date: July 2013

# Worksheet #1 - Approval Page

(UFP-QAPP Manual Section 2.1) Document Title:

Sampling and Analysis Plan, (Field Sampling Plan and Quality Assurance Project Plan), February 2011, Data Gaps Assessment, Tank Farm 2, Category 1 Areas, Naval

Station Newport, Newport, Rhode Island Lead Organization: Naval Facilities Engineering Command Mid-Atlantic Preparer's Name and Organizational Affiliation: Tetra Tech Preparer's Address and Telephone Number: 234 Mall Boulevard Suite 260, King of Prussia. Pennsylvania 19046-1433 Preparation Date (Day/Month/Year): July 2013 Investigative Organization's Project Manager: Signature/Date Dabra Seiken, CG, Tetra Tech Investigative Organization's Project QA Manager: Signature/Date Tom Johnston, PhD, Tetra Tech Lead Organization's Project Manager: Roberto Pagtalunan, PE, NAVFAC Mid Atlantic Lead Organization QA Officer: Signature/Date NAVFAC Chemist, NAVFAC Atlantic Approval Signatures: Signature/Date Kymberlee Keckler, U.S. EPA

Signature/Date

Title: Data Gaps Assessment Document No.: W5211722F Revision Number: 0 Date: July 2013

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Lead Organization: Naval Facilities Engineering Command Mid-Atlantic

Preparer's Name and Organizational Affiliation: Tetra Tech

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Preparation Date (Day/Month/Year): July 2013 Investigative Organization's Project Manager: Signature/Date Dabra Seiken, CG, Tetra Tech Investigative Organization's Project QA Manager: Signature/Date Tom Johnston, PhD, Tetra Tech Lead Organization's Project Manager: Signature/Date Roberto Pagtalunan, PE, NAVFAC Mid Atlantic Lead Organization QA Officer: Signature/Date NAVFAC Chemist, NAVFAC Atlantic KINTSI Approval Signatures: Signat re/Date Kymberiee Keckler, U.S. EPA Signature/Date Pamela Crump, RIDEM

Title: Data Gaps Assessment Document No.: W5211722F Revision Number: 0 Date: July 2013

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	Signature/Date Tom Johnston, PhD, Tetra Tech
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Lead Organization QA Officer:	Signature/Date NAVFAC Chemist, NAVFAC Atlantic
Approval Signatures:	Signature/Date Kymberlee Keckler, U.S. EPA
	Poece

Signature/Date

Pamela Crump, RIDEM

Sité Location: Portsmouth, Rhode Island

Title: Data Gaps Assessment Document No.: W5211722F Revision Number: 0 Date: July 2013

**EXECUTIVE SUMMARY** 

This Sampling and Analysis Plan presents the methodologies to be used for collecting data that will be

used to determine the nature and extent of contamination related to past activities that have resulted in

what are considered releases under the Comprehensive Environmental Response, Compensation and

Liability Act (CERCLA) at Tank Farm 2 (Site 10) which is part of Naval Station (NAVSTA) Newport,

formerly the Naval Education and Training Center (NETC) Newport. The data collected is also expected

to be used to estimate whether risks from exposure of human and ecological receptors to site

contaminants merit actions to further investigate or mitigate the risks, in an effort to protect human health

and the environment.

The NAVSTA Newport is located in the Towns of Newport, Middletown, and Portsmouth, Rhode Island,

approximately 25 miles southeast of Providence. Tank Farm 2 (the Site) is situated at the northern and

central portions of NETC-Newport, in Portsmouth, Rhode Island. The Site is located approximately 1,000

to 1,500 feet east of Narragansett Bay (Figure 1).

The Site occupies approximately 70 acres of land bordered by undeveloped woodlands to the west; Tank

Farm 1, a campground, and a recreational area are located to the northeast; residential housing is

located to the southeast; and Newport Naval Cable TV property and farmland about the site to the south.

The Site is occupied by eleven 2.5-million gallon capacity concrete underground storage tanks (USTs)

(Tanks 19 through 29). Underground petroleum distribution lines connect the USTs to the Naval Fuel

Loading Area (about 1,000-feet northwest of the Site). The Site is covered with overgrown areas

(formerly grass), paved access roads, and tank access chambers. The Site also contains support

buildings, including Building 219, a former transformer building.

The tanks were constructed in the 1940s. The tanks stored No.5 fuel oil from the 1940s to 1975, distillate

fuel from 1975 to 1985, and marine diesel fuel from 1985 until the mid-1990s. Tank 22 was taken out of

service and cleaned in the 1970s and then used as a storage tank for sludge. Tank Farm 2 was operated

by the Navy until 1974, when the property was leased to the Defense Energy Support Center (DESC).

The DESC actively operated the Site until the 1990s, when the tanks were emptied and cleaned. The

DESC still maintains contractual control of the property, although it is not in active operation.

Historical information suggests that sludge from tank cleaning was disposed of on the ground surface in

the vicinity of each tank, from the 1940s to the mid-1970s (Envirodyne Engineers, Inc., 1983). Since that

time, the sludge has reportedly been disposed of at off-site facilities.

Environmental investigations and remediation have previously occurred at the Site by consultants and

contractors hired by the DESC, and have been performed under the Rhode Island Department of

Environmental Management (RIDEM) regulations. The investigations resulted in the installation of 28

monitoring wells, water level and non-aqueous phase liquid (NAPL) gauging rounds, and the collection of

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soil and groundwater samples. Monitoring wells and sampling sites were located near the tanks and in

areas identified in historic aerial photographs as areas of concern. The tanks and portions of the

distribution piping have also been cleaned and inspected, although they have not been permanently decommissioned or given official closure by RIDEM. Soil excavations have occurred at the site to remove

soils determined to be above applicable RIDEM standards based on investigations conducted.

In accordance with decisions made by the Project Team, the Site has been segregated into Category 1

(CERCLA-regulated) areas and Category 2 (RIDEM UST Division regulated) areas, based on the

activities and the contaminants suspected to have been released in each areas. Also, RIDEM has

identified several other areas of potential concern for which the scope of investigation has not been

determined. These additional areas of potential concern are currently termed Category 3 areas, and will

be further evaluated to determine if additional investigation of these areas, if any, will be performed as a

Category 1 or Category 2 area.

This Sampling and Analysis Plan (SAP) addresses only Category 1 Areas. Based on a review of historical

environmental investigations and remediation performed at the Site, data gaps for six Category 1 areas

were identified. The Category 1 areas that will be further characterized under this SAP are:

Four areas of concern (AOCs) where it is suspected that sludge was deposited on the ground and

burned (AOC-001, -003, -004 and -005);

A former transformer building, also known as Building 219, and

The former JP-5 soil pile/ buoy storage area.

Data gaps in these Category 1 AOCs have been identified as follows:

Soil samples from the four AOCs where suspected sludge burning operations took place were not

analyzed for potential site-related contaminants other than petroleum; therefore, additional soil samples

will be collected and analyzed for metals, polynuclear aromatic hydrocarbons (PAHs) and dioxins/furans

in these four Category 1 AOCs. This analyte list includes potential by-products of the burning of

petroleum sludge. Also, at the request of RIDEM, extractable total petroleum hydrocarbon (ExTPH) and

gasoline range organics (GRO) will be analyzed.

Building 219 is a former transformer building. Based on the use of this area for electrical equipment and

the possible presence of PCBs in transformer oil, and previous environmental investigations in this area

which indicated the presence of PCBs in soil, the potential contaminants in this area are PCBs. Data

gaps for Building 219 include adequate definition of the extent of the PCB contamination associated with

the Building and its equipment. Additional sampling and analysis will be performed in this area to further

define the extent of PCB contamination in soil.

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The former buoy storage area is located west of Tank 28. The area was previously investigated as:

AOCs -022, -026, -033, -034, -035 and -036. Soil and groundwater samples in this area have been

collected and analyzed for organic compounds including TPH, VOCs, and SVOCs. Groundwater from

this area has also previously been analyzed for lead. However, soil has not been analyzed for lead.

Therefore, additional sampling and analysis will be performed in this area to determine if a release of lead

has occurred from the reported storage of buoys in this area.

Soil samples at all the investigation areas addressed under this SAP will be collected using a drill rig or

direct-push methods, at depths of 0 to 1, 2 to 4 and 8 to 10 feet, barring areas where shallow refusal does

not allow soil samples collected at all depths. Groundwater samples will not be collected in these areas

because groundwater has been previously sampled and monitored, and results did not suggest

contamination migration from soil to groundwater.

Following completion of the investigations, the Navy will prepare a Data Gaps Assessment (DGA) Report

for these Category 1 areas of the Site. The report will fill the requirement for either a Study Area

Screening Evaluation (SASE) or a Remedial Investigation (RI) Report. This document will summarize the

investigation activities, describe any issues encountered in the field and corrective actions taken, provide

tables comparing soil and groundwater sampling results to screening levels and other applicable criteria,

and provide figures depicting the locations sampled and the spatial distribution of contaminants. The

Data Gaps Report will also contain recommendations for next steps, as necessary. The Draft Data Gaps

Report will be submitted to RIDEM and USEPA for review and approval.

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# **NUMBER**

17-1 Number of Sample Locations by Analytical Group, Matrix, Tank Farm Area, and Depth (soils)

# **FIGURES**

## **NUMBER**

- Site Locus Tank Farm 2
- Site Plan and Groundwater Contour Plan Tank Farm 2

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# FIGURES (cont.)

# **NUMBER**

- 3 Northern Portion of Site, Category 1 AOCs
- First Tier Conceptual Site Model 4
- Proposed Building 219 Sample Locations 5
- Planned Soil Sampling Locations Category 1 / AOC-001 6
- Planned Soil Sampling Locations Category 1 / AOC-003 7
- Planned Soil Sampling Locations Category 1 / AOCs-004 and 005 8
- Planned Former Buoy Storage Area Sample Locations

## **REFERENCES**

### **APPENDICES**

- Α Historic Site Conditions
- Tetra Tech and EPA SOPs В
- С Field Documentation Forms
- D Project-Specific Field Task Procedures
- Ε **Analytical Specifications**
- Laboratory Certification and SOPs

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### **Acronyms**

AOC Area of Concern bgs **Below Ground Surface** °C **Degrees Celsius** 

CCC Calibration Check Compound **CCV** Continuing Calibration Verification

**CERCLA** Comprehensive Environmental Response Compensation and Liability Act

CFR Code of Federal Regulations

Comprehensive Long-Term Environmental Action Navy CLEAN

Coefficient of Determination COD CSM Conceptual Site Model CTO Contract Task Order Direct Exposure Criteria DEC

DESC **Defense Energy Support Center** DFTPP Decafluorotriphenyl-phosphine

**Detection Limit** DL DO Dissolved Oxygen Department of Defense DoD DPT **Direct Push Technology** Data Quality Objective DQO Diesel Range Organic DRO **Data Validation Manager** DVM

EΑ EA Engineering Science and Technology

**Electron Capture Detector** ECD EDB 1.2-Dibromoethane

Electronic Data Deliverable EDD EDL **Estimated Detection Limit** 

**ELAP Environmental Laboratory Accreditation Program** EPA United States Environmental Protection Agency Extractable Total Petroleum Hydrocarbon ExTPH

FID Flame Ionization Detector **FOL** Field Operations Leader

aram

ĞC Gas Chromatograph GRO Gasoline Range Organics Geographic Information System GIS

**HASP** Health and Safety Plan HCL Hydrochloric Acid

HSM Health and Safety Manager IAS Initial Assessment Study

**Initial Calibration ICAL** 

I/C DEC Industrial/Commercial Direct Exposure Criteria

Interference Check Sample ICS **IDW Investigation Derived Waste** Installation Restoration IR IS Internal Standard

Liter

LCS Laboratory Control Sample **LNAPL** Light Non-Aqueous Phase Liquid

LOD Limits of Detection LOQ Limits of Quantitation

MCL Maximum Contaminant Level Milligrams per Kilogram mg/kg

Milliliters mL

**MPC** Measurement Performance Criteria

MS Mass Spectrometer

MS Matrix Spike

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**MSD** Matrix Spike Duplicate NAD North American Datum NAPL Non-Aqueous Phase Liquid

**NAVFAC** Naval Facilities Engineering Command

NAVSTA **Naval Station Newport** 

NETC Naval Education and Training Center

NIRIS Navy Installation Restoration Information System

ORP Oxygen Reduction Potential

OSHA Occupational Safety and Health Administration

**Ounces** 07

Percent Relative Standard Deviation %RSD

%R Percent Recovery

Polynuclear Aromatic Hydrocarbon PAH

Polychlorinated Biphenyl PCB PID Photo Ionization Detector

PM **Project Manager** 

PPE Personal Protective Equipment

parts per million Ppm

**Project Quantitation Limit** PQL **PQO Project Quality Objective** PSL Project Screening Level QΑ **Quality Assurance** 

QAM **Quality Assurance Manager** 

QC **Quality Control** 

QSM Quality Systems Manual

Res DEC Residential Direct Exposure Criteria

RF Response Factor Remedial Investigation RΙ

RIDEM Rhode Island Department of Environmental Management Rhode Island Pollution Discharge Elimination System RIPDES

Relative Percent Difference RPD Remedial Project Manager RPM Relative Retention Time RRT Relative Standard Deviation RSD Regional Screening Level RSL

Retention Time RT

SAP Sampling Analysis Plan

SASE Study Area Screening Evaluation

Sample Delivery Group SDG SIM Select Ion Monitoring

Site Investigation and Remedial Action Report SIRAR

Standard Operating Procedure SOP

**SPCC** System Performance Check Compound

Structured Query Language SQL

SSL Soil Screening Level SSO Site Safety Officer

**SVOC** Semi-Volatile Organic Compound

TBD To Be Determined

Total Chlorinated Volatile Organic Compound tCVOC

TEQ **Toxicity Equivalent** 

TPH Total Petroleum Hydrocarbon

Tetra Tech EC **TtEC** 

UCL Upper Confidence Limit

**USEPA** U.S. Environmental Protection Agency

UST Underground Storage Tank VOC Volatile Organic Compound

### SAP Worksheet #2 -- SAP Identifying Information

Site Name/Number: Tank Farm 2, NAVSTA Newport

**Operable Unit:** 

**Contractor Name:** Tetra Tech, Inc. (Tetra Tech)

**Contract Number:** N62470-08-D-1001

Naval Facilities Engineering Command (NAVFAC) Mid-Atlantic **Contract Title:** 

Comprehensive Long-Term Environmental Action Navy (CLEAN)

Work Assignment Number (optional): CTO WE30

- 1. This SAP was prepared in accordance with the requirements of the Uniform Federal Policy for Quality Assurance Plans (UFP-QAPP) (U.S. EPA 2005) and EPA Guidance for Quality Assurance Project Plans, EPA QA/G-5, QAMS (U.S. EPA 2002).
- 2. Identify regulatory program: Comprehensive Environmental Response, Compensation, and Liability Act of 1980 (CERCLA).
- 3. This SAP is a project-specific SAP.
- 4. List dates of scoping sessions that were held:

	Date	
Introductory Session - Tetra Tech, & Mid Atlantic	10/21/2010	
Introductory Session (RPM Meeting) - Tetra Tech, Mid		
Atlantic, USEPA Region I, RIDEM	11/17/2010	
Technical Session (Tetra Tech)	12/21/2010	

5. List dates and titles of any SAP documents written for previous site work that are relevant to the current investigation.

Little	Date
Draft Work Plan for Site Closure, Tank Farm 2, Foster Wheeler Environmental Corporation.	September 2003
Draft Condensed Work Plan for Soil and Groundwater Sampling, Tank Farm 2,	
Tetra Tech EC, Inc.	May 2005
Work Plan for Monitoring Well Installation, Tank Farm 2, Tetra Tech EC, Inc.	July 2005
Email titled: 'Tank Farm 2 Summary of issues for SAP' (Table A-1 attached).	
Email sent from Tetra Tech to EPA and RIDEM on date indicated.	December 14, 2010

6. List organizational partners (stakeholders) and connection with lead organization:

U.S. EPA, Regulatory Oversight	NAVFAC, Mid-Atlantic – Responsible Party
RIDEM, Regulatory Oversight	Tetra Tech, Contractor to NAVFAC
NAVSTA, Property Holder	

7. Lead organization

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U.S. Navy (NAVFAC Mid Atlantic)

8. If any required SAP elements or required information are not applicable to the project or are provided elsewhere, then note the omitted SAP elements and provide an explanation for their exclusion below:

None		

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# **SAP Worksheet #3 -- Distribution List**

(UFP-QAPP Manual Section 2.3.1)

Name of SAP Recipients	Title/Role	Organization	Telephone Number (Optional)	E-mail Address or Mailing Address	Document Control Number (Optional)
Roberto Pagtalunan, PE	Remedial Project Manager (RPM)	NAVFAC Mid Atlantic	757-341-2010	roberto.pagtalunan@navy.mil	Not applicable (NA)
Darlene Ward	IR Program Contact	NAVFAC Newport	401-841-6376	darlene.ward@navy.mil	NA
TBD	Project Chemist	NAVFAC Atlantic	TBD	TBD	NA
Dave Barclift	Ecological Risk Assessor	NAVFAC	215-897-4913	david.barclift@navy.mil	NA
Kymberlee Keckler	Remedial Project Manager (RPM)	USEPA Region 1 Federal Facilities	617-918-1385	keckler.kymberlee@epa.gov	NA
Pamela Crump	Remedial Project Manager (RPM)	RIDEM Div Site Remediation	401-222-2797	pamela.crump@dem.ri.gov	NA
Dabra Seiken	Project Manager (PM)	Tetra Tech	978-474-8400	dabra.seiken@tetratech.com	NA
Tom Johnston (electronic copy)	Quality Assurance Manager (QAM)	Tetra Tech	412-921-8615	tom.johnston@tetratech.com	NA
Matt Soltis (electronic copy)	Health and Safety Manager (HSM)	Tetra Tech	412-921-8912	matt.soltis@tetratech.com	NA
Kayleen Jalkut	Field Operations Leader (FOL)/Project Geologist/ Site Safety Officer (SSO)	Tetra Tech	978-474-8400	kayleen.jalkut@ tetratech.com	NA
Kelly Carper	Project Chemist	Tetra Tech	412-921-7090	kelly.carper@tetratech.com	NA
Jennifer Obrin	Laboratory PM	Katahdin Analytical Services (Katahdin)	207-874-2400	jobrin@katahdinlab.com	NA
Jill Kellmann	Laboratory PM	TestAmerica – West Sacramento (TestAmerica)	916-374-4402	6-374-4402 jill.kellmann@testamericainc.com	
Glenn Wagner	Administrative Record Manager	Tetra Tech	412-320-2211	glenn.wagner@tetratech.com	NA

Project-Specific Sampling and Analysis Plan Site Name: Tank Farms 2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island

Title: Sampling and Analysis Plan Document No.: W5211722F Revision Number: 0 Date: July 2013

# **SAP Worksheet #4 -- Project Personnel Sign-Off Sheet**

(UFP-QAPP Manual Section 2.3.2)

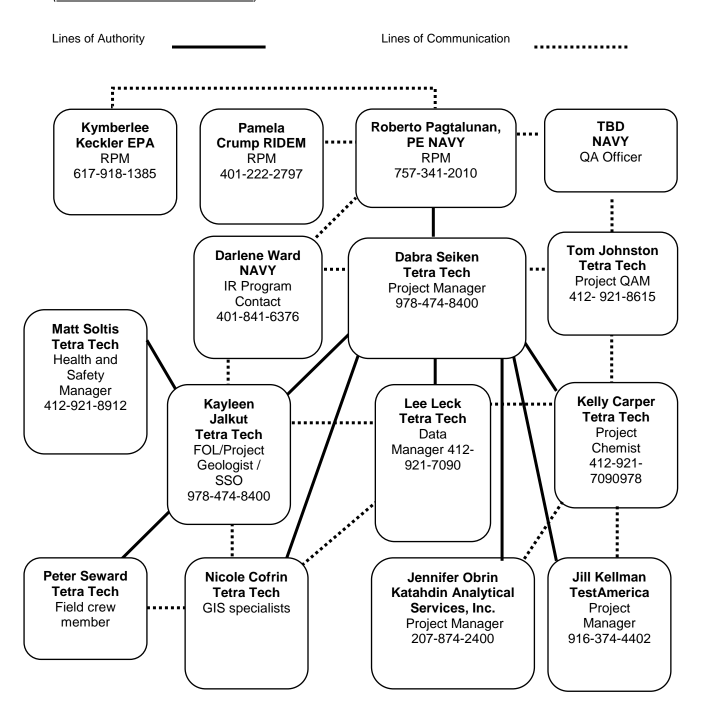
Project personnel who are responsible for implementing portions of the SAP will be provided copies of the applicable SAP sections. Their signatures or email receipt date will indicate that they have read the applicable SAP sections and will perform the tasks as described. If only a portion of the SAP was reviewed, then personnel should note which sections were reviewed.

Name	Organization/Title/Role	Telephone Number (optional)	Signature/email receipt	SAP Section Reviewed	Date SAP Read
Dabra Seiken	Tetra Tech/PM, general project management	978-474-8400	See Worksheet #1	All	
Kayleen Jalkut	Tetra Tech/FOL/Project Geologist, and SSO	978-474-8400		All	
Tom Johnston	Tetra Tech/QAM, quality assurance management, data quality review oversight	412-921-8615	See Worksheet #1	All	
Kelly Carper	Tetra Tech/Project Chemist, laboratory procurement oversight, data quality review, and chemistry support	412-921-7090		All	
Peter Seward	Tetra Tech/Field Sample Collection Specialist/ Sample collection, shipment	978-474-8400		All	
Gary Glennon	Tetra Tech/Database specialist/Geographic Information System (GIS) and analytical data presentation and analysis	978-474-8400		All	
Jennifer Obrin	Katahdin/PM	207-874-2400		Worksheets #15, 19, 20, 23, 24, 25, 28, 30, 34	
Jill Kellmann	TestAmerica/PM	916-374-4402		Worksheets #15, 19, 20, 23, 24, 25, 28, 30, 34	
David Barclift	Navy/ Risk Assessor	215-897-4913		All	

Site Name: Tank Farm2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island Title: Sampling and Analysis Plan Document No.: W5211722F Revision Number: 0 Date: July 2013

# SAP Worksheet #5 -- Project Organizational Chart

(UFP-QAPP Manual Section 2.4.1)



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# **SAP Worksheet #6 -- Communication Pathways**

(UFP-QAPP Manual Section 2.4.2)

Communication Drivers	Responsible Affiliation	Name	Phone Number and/or e-mail	Procedure
Regulatory Agency Interface	Tetra Tech PM	Dabra Seiken	978-474-8400	PM will notify the EPA and RIDEM RPMs at least 48 hours prior to commencement of field activities and 24 hours prior to a change in schedule. PM will provide regulators with weekly field updates via email, including activities performed that week and a schedule of planned activities for the following week. PM will notify regulators via e-mail within 48 hours after receipt of a signed concurrence letter from the Navy RPM to change the scope of work, and prior to execution of the work.
SAP amendments	Navy Remedial Project Manager (RPM)	Roberto Pagtalunan	757-341-2010	RPM sends scope change within 1 week of recognizing need for SAP amendment to Tetra Tech Program office prior to implementing any changes in scope.
Changes in schedule	Tetra Tech PM	Dabra Seiken	978-474-8400	FOL informs PM by phone within same day of recognizing need for change; PM informs RPM by phone within 24 hours and prepares schedule concurrence letter, if deemed necessary by the RPM and PM.
Issues in the field that result in changes in scope of field work	Tetra Tech FOL	Kayleen Jalkut	978-474-8400	FOL informs Tetra Tech PM by phone within same day of identifying field issue. PM approves change same day, if warranted. Document via FMR form.
Issues in the field that result in changes in scope of work	Tetra Tech FOL Tetra Tech PM	Kayleen Jalkut Dabra Seiken	978-474-8400 978-474-8400	FOL informs PM by phone within same day of identifying issue; PM informs RPM by phone within 24 hours, if warranted. PM sends a concurrence letter to Navy RPM, if warranted, within 7 days. RPM signs the letter within 5 days of receipt. Scope change is to be implemented before work is executed. Document the change on a FMR form.
Recommendations to stop work and initiate work upon corrective action	Tetra Tech FOL/SSO Tetra Tech PM Tetra Tech QAM Navy RPM	Kayleen Jalkut Dabra Seiken Tom Johnston Roberto Pagtalunan	978-474-8400 978-474-8400 412-921-8615 757-341-2010	Responsible Party informs subcontractors, the Navy, and Project Team by phone within 1 business day of identifying need to stop work.

Project-Specific Sampling and Analysis Plan Site Name: Tank Farm 2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island

Title: Sampling and Analysis Plan Document No.: W5211722F Revision Number: 0 Date: July 2013

Communication Drivers	Responsible Affiliation	Name	Phone Number and/or e-mail	Procedure
Analytical data quality issues	Katahdin PM TestAmerica PM Tetra Tech Project Chemist Navy RPM	Jennifer Obrin Jill Kellmann Kelly Carper Roberto Pagtalunan	207-874-2400 916-374-4402 412-921-7090 757-341-2010	The Laboratory PM will notify (verbally or via e-mail) the Tetra Tech Project Chemist within one business day of when an issue related to laboratory data quality is discovered.  The Tetra Tech Project Chemist will notify (verbally or via e-mail) the data validation manager (DVM) and the Tetra Tech PM within one business day.  Tetra Tech Project Chemist notifies Tetra Tech PM verbally or via e-mail within 48 hrs. of validation completion that a nonroutine and significant laboratory quality deficiency has been detected that could affect this project and/or other projects. The Tetra Tech PM verbally advises the NAVFAC RPM within 24 hours of notification from the project chemist. The NAVFAC RPM takes corrective action that is appropriate for the identified deficiency. Examples of significant laboratory deficiencies include data reported that has a corresponding failed tune or initial calibration verification.  Corrective actions may include a consult with the NAVFAC Navy Chemist.

Note: Telephone notifications to be documented via email.

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# **SAP Worksheet #7 -- Personnel Responsibilities Table**

(UFP-QAPP Manual Section 2.4.3)

Name	Title/Role	Organizational Affiliation	Responsibilities
Roberto Pagtalunan	RPM	Navy, NAFAC Mid Atlantic	Oversees project implementation, including contract management. Scoping, data review, and evaluation.
Kymberlee Keckler	EPA RPM	USEPA Region I	Participates in scoping, data review, evaluation, and review of the SAP Addendum. Oversees project execution for USEPA.
Pamela Crump	RIDEM RPM	RIDEM, Division of Site Remediation	Participates in scoping, data review, evaluation, and review of the SAP Addendum. Oversees project execution for RIDEM.
Dabra Seiken	PM, Lead Hydrogeologist	Tetra Tech	As PM, oversees project, financial, schedule, and technical day to day management of the project. Provides technical review of interpreted data.  As Lead Hydrogeologist, supervises field work and preparation of geological interpretation and text.
Kayleen Jalkut	FOL/Project Geologist/SSO	Tetra Tech	As FOL, supervises, coordinates, and performs field sampling activities.  As Project Geologist, assimilates geological data, prepares geological interpretation and text.  As SSO, is responsible for staff training and monitoring site conditions related to personnel safety.  Details of the SSO's responsibilities are presented in the site-specific Health and Safety Plan (HASP).
Tom Johnston	QAM	Tetra Tech	Ensures quality aspects of the CLEAN program are implemented, documented, and maintained.
Matt Soltis	HSM	Tetra Tech	Oversees Tetra Tech CLEAN Program Health and Safety Program.
Kelly Carper	Project Chemist	Tetra Tech	Participates in project scoping, prepares laboratory scopes of work, and coordinates laboratory- related functions with laboratory. Oversees data quality reviews and quality assurance of data validation deliverables.
Jennifer Obrin	Laboratory PM	Katahdin	Coordinates analyses with laboratory chemists, ensures that scope of work is followed, provides
Jill Kellmann	Laboratory PM	Test America	quality assurance (QA) of data packages, and communicates with Tetra Tech project staff.
Gary Glennon	Data Manager	Tetra Tech	Consolidates data in database. Analyzes and presents analytical data. Maps or oversees mapping of data in GIS or other system.
Peter Seward	Staff Scientist	Tetra Tech	Collect, package, and ship samples in accordance with the SAP. Assimilates analytical data and prepares text regarding nature and extent of contamination.
Nicole Cofrin	GIS Specialist	Tetra Tech	Map data in GIS.

**Project-Specific Sampling and Analysis Plan** Site Name: Tank Farm 2, NAVSTA Newport

Site Name: Tank Farm 2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island

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SAP Worksheet #8 -- Special Personnel Training Requirements Table

(UFP-QAPP Manual Section 2.4.4)

All field personnel will have appropriate training to conduct the field activities to which they are assigned. Additionally, each site worker will be required to have completed a 40-hour course (and 8-hour refresher, if applicable) in Health and Safety Training as described under Occupational Safety and Health Administration (OSHA) 29 Code of Federal Regulations (CFR) 1910.120(b)(4). Safety requirements are addressed in greater detail in the accompanying site-specific Tetra Tech Health and Safety Plan (HASP), prepared under separate cover.

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## SAP Worksheet #9 -- Project Scoping Session Participants Sheet (UFP-QAPP Manual Section 2.5.1)

Project Name: Data Gaps

Assessment

Projected Date(s) of Sampling: Jan/Feb 2013

(Category 1)

Project Manager: Dabra

Seiken

Site Name: Tank Farm 2, NAVSTA Newport

Site Location: Portsmouth, Rhode Island

Date of Session: October 21, 2010

Scoping Session Purpose: Develop project quality objectives using EPA DQO process

Name	Title	Affiliation	Phone #	E-mail Address	Project Role	
Roberto Pagtalunan	RPM	NAVFAC Mid- Atlantic	757- 341- 2010	roberto.pagtalunan@navy.mil	Remedial Project Manager	
Winoma Johnson	RPM	NAVFAC Mid- Atlantic	757- 341- 2008	Winoma.johnson@navy.mil	RPM and Team Leader	
Stephen Parker	Project Manager	Tetra Tech	978- 474- 8400	Stephen.Parker@tetratech.co m	Facility coordinator	

Comments/Decisions:

Agreed to approach site using same methodology as Tank Farm 4/5 sites, where hazardous materials releases are addressed as Category 1 under CERCLA, and fuel / petroleum and related releases are addressed as Category 2 under RIDEM UST regulations. At this site AOCs where uncontrolled burning of sludge is suspected will

be Category 1 areas.

Use of Category 3 is uncertain.

Action Items: None

Consensus Decisions: None **Project-Specific Sampling and Analysis Plan** Site Name: Tank Farm 2, NAVSTA Newport

Site Name: Tank Farm 2, NAVSTA Newpo Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island Title: Sampling and Analysis Plan Document No.: W5211722F Revision Number: 0 Date: July 2013

Project Name: Remedial

Investigation

Projected Date(s) of Sampling: Jan/Feb 2013 (Category 1)

Site Name: Tank Farm 2, NAVSTA Newport

Site Location: Portsmouth, Rhode Island

Project Manager: Dabra Seiken

Date of Session: November 17, 2010

Scoping Session Purpose: Develop project quality objectives using EPA DQO process

Name	Title	Affiliation	Phone #	E-mail Address	Project Role
Dabra Seiken	Project Manager	Tetra Tech	978-474- 8400	dabra.seiken@tetratech.com	Project manager, Lead Geologist
Roberto Pagtalunan	RPM	NAVFAC Mid-Atlantic	757-341- 2010	roberto.pagtalunan@navy.mil	Remedial Project Manager
Kymberlee Keckler	RPM	USEPA	617-918- 1385	kymberlee.keckler@epa.gov	Remedial Project Manager
Stephen Parker	Project Manager	Tetra Tech	978-474- 8400	stephen.parker@tetratech.com	Facility coordinator
Gary Jablonski	RPM	RIDEM	401-222- 2797	Gary.Jablonski@dem.ri.gov	Remedial Project Manager

Comments/Decisions: For Tank Farms 4 and 5, Category 3 areas are areas that do not fall into Category 1 or 2. A shed identified by RIDEM was the only category 3 area. However, at Tank

or 2. A shed identified by RIDEM was the only category 3 area. However, at Tank Farm 2, there are PCBs in soil near transformers. If facilities are operational (i.e. power is active to a transformer), that would be a facility issue and releases would be reported to RIDEM and likely addressed under state jurisdiction by the facility directly, and not through the IR program, and therefore not be addressed under CERCLA.

The Navy concluded that existing AOCs associated with active utilities will not be

addressed under this remedial investigation.

Action Items: None

Consensus Decisions: None

**Project-Specific Sampling and Analysis Plan** Site Name: Tank Farm 2, NAVSTA Newport

Site Name: Tank Farm 2, NAVSTA Newpo Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island Title: Sampling and Analysis Plan Document No.: W5211722F Revision Number: 0 Date: July 2013

Project Name: Data Gaps

Investigation

Projected Date(s) of Sampling: Jan/ Feb 2013 (Category 1)

Site Name: Tank Farm 2, NAVSTA Newport

Site Location: Portsmouth, Rhode Island

Project Manager: Dabra

Seiken

Date of Session: December 21, 2010

Scoping Session Purpose: Develop project quality objectives

Name	Title	Affiliation	Phone #	E-mail Address	Project Role		
Dabra Seiken	Project Manager	Tetra Tech	978-474- 8400	dabra.seiken@tetratech.com	Project manager, Lead Geologist		
Ann Franke	Chemist	Tetra Tech	978-474- 8400	ann.franke@tetratech.com	Chemistry support		
Stephen Parker	Project Manager	Tetra Tech	978-474- 8400	stephen.parker@tetratech.com	Facility coordinator		
Amy Carey	Geologist	Tetra Tech	978-474- 8400	amy.carey@tetratech.com	Field Staff		

Comments/Decisions: Discussed technical details of sampling program.

The team agreed to conduct investigations at the four former sludge burning pit AOCs, placing borings within each area. Borings will be located in a square (15 foot by 15 foot) grid pattern across each area to allow delineation of the extent of contamination and collect data to represent the conditions in the surface and subsurface soil within the area.

The study areas will be established to be 50% larger than the sludge burning area. The four units range in size from 600 square feet to 4,800 square feet. This limit is intended to reflect a conservative exposure area for an industrial worker, where that worker's duties may keep him/her exposed to the impacted soil.

Action Items: None

Consensus Decisions: None

Site Name: Tank Farm 2, NAVSTA Newport

Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island

SAP Worksheet #10 -- Conceptual Site Model

(UFP-QAPP Manual Section 2.5.2)

10.1 SITE LOCATION AND BACKGROUND

Tank Farm 2 ("the Site") is located in the Melville section of Portsmouth, Rhode Island, just north of

Newport, Rhode Island, close to the eastern shore of Narragansett Bay (Figure 1). It is situated on the

crest of a hill, and the topography across the property slopes westward toward the bay. Melville North

pond is the closest surface (fresh) water body, located approximately 400 feet northeast of the Site

(Figure 1). Groundwater flow at the Site is to the west and northwest.

The Site encompasses approximately 70 acres. It is bordered by undeveloped woodlands to the west

and the Naval Fire Department to the northwest. The Melville Campground and Recreational Area

borders the Site to the north and east. The Newport Naval Cable TV property and the Melville Naval

Family Housing complex are located to the southeast and south (Figure 2).

The ground surface of the Site is currently covered in vegetation, such as brush and grasses, with a few

clear areas along paved access roads. The property is enclosed along the perimeter with a security

fence. Access to the property is via Defense Highway, which runs along the western border of the Site,

between the property and Narragansett Bay. The Site has eleven 2.5-million gallon capacity, concrete,

USTs (Tanks 19 through 29) (Figure 2), that are installed in blasted bedrock sockets.

The Site also contains:

1) underground fuel distribution lines installed 10-feet below grade in concrete-lined utility trenches;

2) an underdrain system (ring drain) around each UST that collects and transfers excess

groundwater away from the USTs;

3) sump pump chambers in the pump house next to each UST that contain pumps for the fuel

distribution lines and the ring drain;

4) buried piping connecting the fuel distribution lines and the pump houses;

5) a UST vent and a gauging house connected to each UST; and

6) support buildings, including Building 219 (B219), a former electrical service/transformer building.

In accordance with decisions made by the project team (Worksheet #9), the AOCs at the Site have been

separated into Category 1 (CERCLA- regulated), Category 2 (RIDEM UST Division regulated) areas, and

Category 3 areas (regulatory pathway not yet defined). This SAP has been prepared to address the

Category 1 areas only.

The categories of areas within the tank farms are defined as follows:

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1. Category 1 (CERCLA-regulated) areas - these are locations of releases/suspected releases of

CERCLA hazardous substance(s), which were not the result of DESC's petroleum operations.

2. Category 2 (RIDEM UST Division regulated) areas – these locations relate to petroleum contamination resulting from DESC operations, and which have been, or are currently being,

addressed by DESC.

3. Category 3 areas (additional areas of concern) – areas for which the scope of investigation has

not been determined, a release is not confirmed, and a regulatory pathway is not yet defined.

The Category 1 areas consist of three types of areas/contaminants at Tank Farm 2:

1) Contaminants associated with burning of tank sludge (polynuclear aromatic hydrocarbons

[PAHs], metals, and dioxins. Sludge may have been deposited on the ground in areas of the

Site. There is some evidence that in some of these areas, the sludge was burned. Burning

sludge can alter the petroleum sludge, potentially producing dioxins and pyrogenic PAHs,

and release elevated concentrations of heavy metals, all of which may have deposited onto

soil. Heavier petrogenic PAHs that are not combustion related products may also have been

released by this process to the surface soil. The USEPA has stated that areas where

evidence of sludge burning has taken place would be governed by CERCLA.

2) Areas where polychlorinated biphenyls (PCBs) may have been released to the environment.

The USEPA has stated that the areas where PCB- containing oils were used would be

governed by CERCLA.

3) Areas where lead may have been released to the environment from the storage of painted

buoys. The USEPA has stated that areas where lead-based paint has been released to the

environment could be governed by CERCLA.

The Category 1 portions of the Site that have not been adequately characterized with respect to PAHs,

metals, dioxins and/or PCBs and require further investigation are:

1) AOC 001, AOC-003; AOC-004 and AOC-005. (Areas where sludge may have been

deposited on the ground and burned);

2) Building 219 (Former Transformer Building), and

3) Former buoy storage area.

10.2 SITE HISTORY

The US Navy has owned the Site since at least the 1940s. The USTs were constructed in the 1940s. The

tanks stored No.5 fuel oil from the 1940s to 1975, distillate fuel from 1975 to 1985, and marine diesel fuel

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from 1985 until the mid-1990s. Tank 22 was taken out of service and cleaned in the 1970s and then used

as a storage tank for sludge. The tanks still remain on Site although they have not been used for fuel

storage since tank closure activities were performed in 1996 and 1997. The Site was operated by the

Navy until 1974, when the property was leased to the DESC. The DESC actively operated the Site until

the 1990s, when the tanks were emptied and cleaned. The DESC still maintains contractual control of

the property although it is not in active operation.

The USTs at the Site were periodically cleaned and the bottoms of the tank pumped to remove

accumulated sediments and water (sludge). Historical information suggests that this sludge was released

to the surface soil in the vicinity of each tank, from the 1940s to the mid-1970s (Envirodyne Engineers,

Inc., 1983). Since that time, the sludge was reportedly disposed of at off-site facilities. In addition, sludge

was reportedly stored in Tank 22. The open burning of sludge in four locations (AOC-001; -003; -004,

and -005) near Tanks 19 and 21 (Figure 3) was postulated, based on an analysis of aerial photographs

from the 1940s to the 1990s (TtEC, 2005).

All of the tanks and accessible tank equipment (pumps, interior pipelines and vaults) were emptied and

cleaned in 1996/1997 and have not been used for fuel storage/distribution since that time. The piping

was decommissioned. The USTs were also structurally inspected after the cleaning and all were found to

contain cracks in the sides or bottoms. Tanks and fuel distribution piping were again cleaned in 2001.

The USTs were ballasted with about 1.25 million gallons of water each. The results of the cleaning and

decommissioning activities are provided in Appendix A, Table A-1.

Environmental investigations and remediation were previously performed at the Site by consultants hired

by the DESC. These investigations were performed under the RIDEM regulations. The DESC and Navy

are working to turn control of the Site back to the Navy.

10.3 **GEOLOGY** 

The Site is located in the southeastern portion of the Narragansett Basin. The basin is underlain by

Pennsylvanian age, non-marine, sedimentary and metamorphic rocks, including the Rhode Island

Formation. Bedrock at the Site is described mostly as shale that is weathered and/or metamorphosed.

Siltstones, sandstones and conglomerates have also been observed at the Site, but in much less

abundance. Bedrock is generally observed 5 to 15 feet below ground surface (bgs). Overburden

materials consist of glacial sediments that are a mixture of silts, sands, and gravels and usually classified

as glacial till. Fill, which is usually reworked material from the Site, is also present in the overburden.

Groundwater is usually encountered approximately 5 to 30 feet bgs. The water table is usually within the

bedrock. Groundwater flows in a westerly/northwesterly direction toward Narragansett Bay.

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10.4 SUMMARY OF ENVIRONMENTAL WORK CONDUCTED

The first environmental investigation was an Initial Assessment Study (IAS) completed by Envirodyne

Engineers Inc. in 1983. Upon completion of the IAS, the Site was recognized as an area that required

further environmental investigation, because petroleum sludge had been placed on the ground, but no

samples were collected from the Site at that time.

The following sections list investigative and remedial actions previously conducted at the Site under the

direction of the DESC.

10.4.1 <u>Tank Closure</u>

Tank closure activities took place between September 1996 and May 1997 under the direction of DESCs

environmental consultant. Additional tank and pipe cleaning was completed by DESCs environmental

consultant between January and August 2001. These closure activities for each of the 11 tanks are

detailed in Appendix A, Table A-1.

10.4.2 <u>Monitoring Well Installation and Sampling</u>

GZA GeoEnvironmental, Inc. (GZA) installed 11 monitoring wells (one adjacent to each UST) between

October and December of 1996 (GZ-201 to GZ-211) and an additional 17 monitoring wells between

September and October 1997 (GZ-212 to GZ-228) (Figure 2). Several sampling events were conducted

from 1997 to 2009, with groundwater samples analyzed at various times for total petroleum hydrocarbons

(TPHs), volatile organic compounds (VOCs), semivolatile organic compounds (SVOCs), PAHs, and total

and dissolved lead. Wells GZ-202 and GZ-208 were sampled once for an oil-water mixture and the lab

performed a petroleum fingerprint analysis on the samples. The hydrocarbons in the samples

represented hydrocarbons in the Number 2 Fuel Oil boiling point range. Details of the groundwater

sampling events are presented in Appendix A, Table A-2. The sample results for the wells associated

with each tank are presented in Appendix A, Tables A-3.1 through A-3.13.

Table A-1 (Appendix A of this SAP) summarizes the monitoring wells, the analytical fractions analyzed,

and the exceedances of RIDEM standards associated with each tank. LNAPL was detected in the most

recent gauging/sampling rounds (2009) in wells GZ-202, GZ-208, GZ-211 (Tanks 20, 26, and 29,

respectively). No other exceedances were found in the groundwater samples.

10.4.3 Soil Boring Sampling

During the installation of monitoring wells in 1996 (GZ-201 to GZ-211) and 1997 (GZ-212 to GZ-228), soil/

weathered bedrock samples were collected every 5 feet from the ground surface to the bottom of the

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boring, and sample headspace was screened for total VOCs using a photo-ionization detector (PID) and

flame-ionization detector (FID). Typically, the soil/weathered bedrock sample from each boring that

demonstrated the highest total VOC concentrations was sent to the laboratory. Twenty-two

soil/weathered bedrock samples were submitted to the laboratory for TPH, VOC and PAH analyses.

In May 1997, GZA completed 35 shallow soil borings (B-1 to B-35) along the fuel distribution lines (Figure

2). Borings were advanced to a depth of approximately 12-feet. One soil sample was collected from

each boring at a depth of approximately 10 to 12 feet, which is just below the 10-foot deep concrete lined

utility trench that houses the fuel distribution lines (Figure 2). The sample was screened using a PID and

a FID and then submitted to the laboratory for TPH analysis.

Soil sample analytical results are summarized in Tables A-3.1 through A-3.13, presented in Appendix A.

As summarized in Table A-1 and in detail on Table A-3.9, in 1996 a soil sample collected at 15 to 17 feet

bgs in boring GZ-209 exceeded the RIDEM GB Leachability Criteria with a TPH concentration of 5,600

milligrams per kilogram (mg/kg). However, the groundwater sample results from well GZ-209 did not

exceed the RIDEM standard. No other exceedances were identified for the soil samples.

10.4.4 **Soil Testing & Excavation** 

From May 2005 to June 2006, Tetra Tech EC (TtEC) performed soil testing and remedial excavations at

the Site in support of a SIRAR. This work was done in accordance with the following work plans:

Draft Work Plan for Site Closure, Tank Farm 2, September 2003, Foster Wheeler Environmental

Corporation.

Draft Condensed Work Plan for Soil and Groundwater Sampling, Tank Farm 2, February 2005,

Tetra Tech EC, Inc.

These work plans identified AOCs to be investigated at the Site. The initial (2003) work plan identified

AOCs based on previous sampling and gauging activities on site. As a result of RIDEM comments, the

subsequent (2005) work plan identified historical AOCs based on the analysis of aerial photographs from

the 1940s to the 1990s. These historical AOCs were explored during the 2005-2006 SIRAR.

A complete list of the AOCs explored appears in the July 2006 SIRAR (TtEC, 2006). AOCs numbered

TF2-001 to TF2-043 were identified based on historical aerial photographs, and the locations of an

additional 14 AOCs (not numbered) were identified by RIDEM and included in the investigation. These

additional AOCs included the areas surrounding the transformers and power poles on the Site, as well as

the soils touching the side walls and below the vents of each tank. Additional information regarding these

AOCs is provided in the SIRAR (TtEC, 2006).

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Soils at the explored areas were tested and analyzed and, based on the results; remedial excavations

were performed, where necessary. These activities are summarized in Appendix A in Table A-1 and

described below.

Soil Testing

The AOCs (identified by aerial photography) (TF2-001 through TF2-043) were evaluated in the field in

December 2004. Nine AOCs (008, 011, 021, 025, 032, 038, 039, 040, and 041) were determined to be

physical features, construction related, or bedrock outcrops that had the appearance of ground staining

on the aerial photos. As a result, these AOCs were not investigated in the work that followed and are no

longer considered AOCs.

The remaining AOCs were investigated during the SIRAR using Petroflag<sup>™</sup> screening for TPH. Shallow

test pits were dug within each AOC and samples were collected from the four sidewalls and the bottom of

the test pits. Test pits ranged in length from 24 to 254 feet, depending on the size of the AOC. Samples

were screened for TPH using Petroflag<sup>™</sup> and the samples with results >100 parts per million (ppm) were

analyzed for TPH by the laboratory.

The laboratory TPH analysis included diesel range organics (DROs) and/or GROs. DRO and GRO

analyses were performed for the AOCs associated with the JP-5 soil piles identified on historic aerial

photographs, these included AOCs 022, 026, 033, 034, 035, and 036. All other samples only had DRO

analysis. The location of the former JP-5 soil pile, also the general area where Naval buoys were

formerly stored, is shown on Figure 2. If laboratory analysis revealed a TPH concentration between 100

and 500 mg/kg, the sample was tested for VOCs and SVOCs. If TPH concentration was greater than

2,500 mg/kg (RIDEMs industrial/ commercial [IC] direct exposure concentration [DEC]), the area was

automatically flagged for remediation (AOC 028, AOC 037, Tank 25 sidewall). Where lab analysis

revealed a TPH concentration of greater than 500 mg/kg (RIDEMs Residential DEC [RDEC]) but less

than 2,500 mg/kg, the sampling location was re-sampled. Re-sampled areas were also analyzed for

VOCs and SVOCs if TPH results were greater than 100 mg/kg.

In addition to the AOCs, each of the eleven tanks at the Site was investigated during the SIRAR by

excavating test pits adjacent to the downgradient side of each tank to a depth of 10 feet and sampling

and analyzing soil from the excavations. One base sample and four sidewall samples were collected at a

depth of 5 feet and one base and four sidewall samples were collected at a depth of 10 feet. Soil

samples were screened with Petroflag<sup>TM</sup> and results greater than 100 parts per million (ppm) were

analyzed for TPH by a laboratory. Within the test pit adjacent to Tank 25, a release was discovered and

remedial action, involving large scale soil (over 2,000 cubic yards) excavation, was performed. Soil was

excavated until confirmatory analysis showed remaining soil contaminant concentrations to be below

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RIDEMs RDECs and ICDECs for VOCs, SVOCs, and TPH (Refer to Tables A-1 and A-3.1 through A-

3.13, Appendix A).

Beneath each tank vent surface soil sampling was conducted. A soil sample from 1-foot depth was

collected beneath each vent and analyzed for TPH. TPH was not detected above RIDEMs ICDEC.

Soil samples were collected at Building 219 during the SIRAR in June of 2005. Four surface soil samples

(0-6 inches) were collected (one at each side of the building) and analyzed for VOCs and PCBs. Aroclor

1260 was detected above RIDEMs ICDEC and RDEC (2 samples) and EPAs industrial RSL (three

samples) and EPAs residential RSL (all samples).

The former buoy storage area was investigated as several AOCs: AOC-022, -026, -033, -034, -035 and -

036 (Table A-1, Appendix A). The results of the soil and groundwater sampling and analysis in this area is

detailed in Appendix A. In summary, results of the sampling and analysis indicated some elevated

petroleum-related constituents, but not at concentrations requiring remedial action.

**Remedial Excavations** 

The DESC performed remedial excavations in areas where contamination was related to fuel storage and

distribution and the TPH concentrations from laboratory analysis of test pit samples were above the I/C

TPH DEC standard. As shown in Appendix A, Table A-1, remediation was performed at Tank 25, AOC-

028, and AOC-037.

Soil was excavated outward from the location of the soil sample that exceeded the I/C TPH DEC

standard. The excavation was directed toward petroleum contaminated soil by TPH screening using

Petroflag<sup>™</sup>, which was performed on soil from the sidewalls and base of the excavations. Soil samples

were taken from the sidewalls and bottoms of the excavations to determine whether removal of

contaminated soil was complete. These samples were sent to the laboratory for analysis. There were no

post-remedial excavation exceedances of the I/C TPH DEC for TPH.

10.5 **CONCEPTUAL SITE MODEL** 

Figure 4 presents the conceptual site model (CSM) for the Site. The CSM is also described below:

Potential contaminants associated with the burning of sludge (PAHs, dioxins, and furans [hereafter

referred to as "dioxins"], and metals) were likely released, by placing combustion sludge on the

ground, at the former burn locations shown on Figure 3. Although fuels formerly stored in the Tanks

contain only trace amounts of heavy metals, these trace concentrations become concentrated in tank

bottom sludges and more concentrated following combustion of the sludge.

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o Small quantities of constituents could dissolve from the sludge and migrate vertically downward,

potentially reaching groundwater. These contaminants could migrate in groundwater

downgradient to the west, which is the direction of groundwater flow at the tank farm.

Groundwater testing in monitoring wells located between 100 and 200 feet downgradient (west)

of these areas indicates that groundwater has not been impacted by sludge burning.

Small quantities of constituents could migrate in the smoke from the combustion of this material.

These constituents would have dispersed and been deposited via aerial deposition. The

concentrations from this deposition mechanism would be diluted due to the large area over which

the deposition would occur.

Small quantities of constituents could also be transported over the ground surface via overland

flow following a precipitation event. However, the vast majority of the constituents are expected

to be confined to the former burn areas and immediately surrounding and beneath the former

burn areas.

PCBs, as Aroclor 1260, associated with the transformer building (Building 219), have been

detected in surface soil. The locations with the highest concentration of PCBs are near the doors

to the building (Figure 5). These are likely associated with the use of PCB-containing transformer

oils in the transformers in the building. The release(s) are associated with incidental spillage to

the surface soil during routine use and maintenance of the transformers in the building.

The storage of Naval buoys was observed on aerial photography in a portion of the site west of Tank

28. Lead-based paint from buoys could have weathered and lead could have been released to the

soil.

Currently, the site is largely unused and exposure to human receptors is limited. There is some limited

hunting allowed by Navy personnel, otherwise the Site is unused. The current receptors to environmental

contamination include terrestrial biota and human receptors that could be exposed to surface soil.

Potential human receptors include trespassers and limited recreational users (hunters). Subsurface soil

is not accessible, as the site is unused and vacant. Groundwater is not accessible, as there are no water

supply wells at the site. Potential terrestrial ecological receptors, such as plants, soil invertebrates,

mammals, birds, and reptiles, can be exposed to contaminated surface soil. Most terrestrial receptors are

not substantially exposed to subsurface soils. There are no surface water bodies or sediment at the Site.

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### 10.6 AREAS REQUIRING FURTHER INVESTIGATION

As described in Section 10.4.4, Site soils were removed by the DESC in areas where contamination was related to fuel storage and distribution, with TPH used as an indicator of contamination. In the course of reviewing the various investigations described above, areas requiring further testing/exploration were A description of these areas is provided in Appendix A, Table A-1. The additional investigations required are presented in the column titled "Next Steps"; all areas with an entry other than "No further action" in that column require further investigation.

Similar to how Tank Farms 3, 4, and 5 are being handled, continued investigation, remedial efforts and site closure at the Site fall under different regulatory categories. This SAP is only for the Category 1 areas that are currently identified:

- AOC TF2-001; former suspected tank sludge burn area (Figure 3)
- AOC TF2-003; former suspected tank sludge burn area (Figure 3)
- AOC TF2-004; former suspected tank sludge burn area (Figure 3)
- AOC TF2-005; former suspected tank sludge burn area (Figure 3)
- Building 219; former Transformer Building (Figure 2)
- Former Naval Buoy storage area (Figure 2)

This SAP addresses the required investigations for these areas, as identified in Appendix A, Table A-1.

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SAP Worksheet #11 -- Project Quality Objectives/Systematic Planning Process Statements (UFP-QAPP Manual Section 2.6.1)

The following text describes the development of project quality objectives (PQOs) using the USEPA data quality objective (DQO) process. The primary data users of this investigation will be Tetra Tech and the

Navy.

11.1 PROBLEM STATEMENTS

Former Sludge Burning Areas - Several AOCs at the Site (AOC TF2-001, AOC TF2-003, AOC TF2-004, and AOC TF2-005) have been identified as having a history of burning of tank bottom sludge (Figure 3). Burning of petroleum sludge can potentially alter the sludge, releasing dioxins, metals, and PAHs to soils at concentrations exceeding risk screening criteria. The nature and extent of such contamination has not been established. The Project Team determined that contaminants related to sludge burning will be classified as Category 1, and an investigation and evaluation of environmental media under Category 1 shall be conducted so that, if necessary, a CERCLA risk assessment can be performed. Therefore, data must be collected in accordance with the Navy and USEPA policies for conducting risk assessments under CERCLA. Additionally, regardless of the category of the AOC, RIDEM has requested that TPH

data be collected at the former sludge burning areas.

In order to determine whether a risk assessment is necessary, the following problem must be resolved:

**Problem:** The Navy must determine the nature and extent of contamination related to burning of sludge in soil at AOC TF2-001, AOC TF2-003, AOC TF2-004, and AOC TF2-005, and must conduct a

risk screening of data collected that represents the current conditions of these areas.

Building 219 – Former Transformer Area – Previous surface soil sampling around the building indicated the presence of PCBs (Aroclor 1260) at concentrations up to 18,000 μg/Kg total PCBs, greater than screening criteria. However, the extent of PCB-contaminated soil must be determined. The Project Team determined that contaminants related to Building 219 will be classified as Category 1, and an investigation and evaluation of environmental media under Category 1 shall be conducted so that, if necessary, a CERCLA risk assessment can be performed. Therefore, data must be collected in accordance with the Navy and USEPA policies for conducting risk assessments under CERCLA. In order

to determine whether a risk assessment is necessary, the following problem must be resolved:

**Problem:** The Navy must determine the extent of contamination related to surface spills of PCB-containing oil at Building 219, and must conduct a risk screening of data collected that represents the current conditions of this area.

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Former Buoy Storage Area -Several AOCs at the Site (AOC -022, -026, -033, -034, -035, and -036)

have been identified for the area where Naval buoys were formerly stored. Buoys are often painted with

lead-based paint that can potentially cause a release of lead to the soil at concentrations exceeding risk

screening criteria. The Project Team determined that contaminants related to lead-based paint will be

classified as Category 1, and an investigation and evaluation of environmental media under Category 1

shall be conducted so that, if necessary, a CERCLA risk assessment can be performed. Therefore, data

must be collected in accordance with the Navy and USEPA policies for conducting risk assessments

under CERCLA. In order to determine whether a risk assessment is necessary, the following problem

must be resolved:

Problem: The Navy must determine the nature and extent of lead contamination, if any, related to

storage of Naval buoys, and must conduct a risk screening of data collected that represents the

current condition of these areas.

11.2 **IDENTIFY INPUTS TO PROBLEM RESOLUTION** 

The inputs needed to resolve the problems identified in Section 11.1 include field measurements,

laboratory chemical data, and PSLs as described below. Field tasks to be performed to collect these data

inputs are summarized in Worksheet #14.

11.2.1 **Laboratory Chemical Data** 

The following Category 1 chemical data from fixed-base laboratory analyses are needed and the list of

target analytes is presented in Worksheets #15a:

For the former sludge burning area problems, concentrations of PAHs, TPH, dioxins, and metals in

surface and subsurface soil are needed. These analytical groups were identified as the most likely

classes of contaminants associated with the burning of petroleum sludge and these data are needed

to determine if a risk assessment is necessary. Category 1 data are not needed for groundwater at

this time. The project team will assess the need for groundwater data after the analytical results from

the soil sampling are received.

For Building 219 (former transformer building) PCBs were identified in surface soil in the vicinity of

Building 219. PCBs are the most likely class of contaminants associated with the release(s) in the

vicinity of Building 219 that has not been adequately characterized. Additional PCB data are also

needed to determine if a risk assessment is necessary.

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For the former buoy storage area, lead is the most likely contaminant that has not been previously

assessed and that can be associated with storage of painted Naval buoys. Lead in soil data is

needed to augment the lead in groundwater data and to determine if a risk assessment is necessary.

There is no surface water or sediment at the Site.

11.2.2 **Project Screening Levels** 

The newly-collected chemical data will be screened against project screening levels (PSLs) to determine

if laboratory quantitation limits were adequate. (However, in order to resolve the project problems and

make decisions, separate screening levels are described in Section 11.4.) For this project, there are

PSLs for surface and subsurface soil. These PSLs are identified on Worksheet #15 and were selected

using the following rationales:

Surface soil PSLs - The PSLs are the lowest of the applicable human health risk-screening criteria

(EPA RSLs for residential and industrial soil; the EPA soil to air SSLs and the RIDEM RDEC), the

RIDEM leachability criteria and the selected ecological soil screening levels (SSLs), for the receptors

identified in Section 10.5.

Subsurface soil PSLs - Ecological risk is only applicable for surface soil. Therefore, the PSLs are

the lowest of the same risk-screening criteria as for surface soil, excluding the ecological SSLs.

(Note: PSLs are subject to change, based on ongoing research, and updated values will be used when

screening is performed. PSLs that are current at the time of the risk screening will be used.)

Fixed laboratory analytical methods must be selected such that the subcontracted laboratories can

achieve limits of quantitation (LOQs) less than or equal to the PSLs, to the extent technically feasible

using conventional methods. To simplify the sampling and analysis procedures, the lowest of the surface

and subsurface PSLs for each analyte was designated as the "soil PSL", and method selection was

based on this lowest value for all of the types of soil. Worksheet #15 present the PSLs; the selected

methods; and the laboratory LOQs, limits of detection (LODs), and detection limits (DLs) for each analyte,

for soil.

The laboratories will measure concentrations of analytes except dioxins down to the laboratory DL, and of

dioxins down to the sample-specific estimated detection limit (EDL). Positive detections of analytes

except dioxins between the LOQ and the DL, and of dioxins between the LOQ and the EDL, will be

qualified as estimated "J". The "J" alerts the data user to the increased uncertainty at concentrations

between the DL and LOQ. Use of J-flagged data to achieve project goals is acceptable; however, greater

scrutiny will be applied to J-qualified data.

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Non-detected results will be qualified as "U" and will be reported with an associated value of the LOD,

except for dioxins. Non-detected dioxins results will be reported with an associated value of the EDL, as

provided in the analytical method.

For the purpose of making the decisions identified in Section 11.4, non-detected results with associated

values greater than the screening level will be treated as values that are less than the screening level if

the chemical was not detected in site media during this investigation or in previous investigations;

otherwise, such results will be assigned a value equal to one-half the LOD (or, for dioxins, one-half the

EDL). The limitations on data usability due to unmet sensitivity goals will be evaluated as described in

Worksheet #37 and will be discussed in the project report. The data usability assessment will consider

uncertainty associated with LOQ and/or LOD and EDL values that are greater than the PSL and will

evaluate whether the inability to detect or quantify an analyte at levels equal to or less than the PSL

creates a data gap that has an adverse effect on decision making.

The background data set for various media at NAVSTA Newport will also be used to determine whether

metals present onsite are naturally occurring or site-related. Background data are described in the

"Basewide Background Study Report for Naval Station Newport, Newport Rhode Island" (Tetra Tech, July

2008).

11.3 STUDY BOUNDARIES

Aerial photographs and data previously collected in conjunction with soil and groundwater sampling and

removal actions were used to determine the areas on which to focus the investigation covered in this

SAP. As described in Worksheet #10, Section 10.8, the areas to be investigated were categorized as to

the appropriate regulatory path.

The areas of focus for Category 1, where soil data potentially will be used for CERCLA-type risk

assessment, are four areas where tank bottom sludge was presumably burned (AOC TF2-001, AOC TF2-

003, AOC TF2-004, and AOC TF2-005 (Figure 3) and one area (Building 219) where PCBs were

released to the ground surface (Figure 5) and one area where Naval buoys were stored (Figure 2).

Extent of AOC boundaries were based upon mapped areas (mapped via aerial photography) of the AOC

as provided in the Site Investigation and Remedial Action Report (TtEC, 2006). Two general soil

populations must be represented in order to resolve the Category 1 problems - soil that potentially

contains chemicals related to sludge-burning, PCB releases, or lead based paint releases at

concentrations that exceed the PSLs and background concentrations, and soil that does not.

The soil depth intervals of interest are those set forth by USEPA policy for risk assessment to define

surface soil and subsurface soil. Surface soil is be defined as soil collected to 1 foot bgs. Subsurface soil

is defined as soil collected between 1 and 10 feet bgs or to top of bedrock, whichever is shallower.

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Contaminants from sludge-burning (AOC-003, -004 and -005), or lead based paint releases (buoy storage

area) do not appear to have impacted groundwater because previous groundwater sampling

downgradient of these areas did not indicate contamination (Appendix A). Furthermore, the project team

has agreed to examine the soil data before making the determination whether groundwater sampling is

warranted. Therefore, no groundwater data are necessary for the Category 1 areas. However, the

screening of soil data against protection of groundwater SSLs is of interest to allow a qualitative

evaluation of the potential for chemical migration from soil to groundwater, in the unlikely event that

contaminants migrated through the bedrock.

11.4 DEVELOP THE ANALYTIC APPROACH

The rules described in this section will be used to evaluate the newly-acquired and usable historical

chemical data and to make decisions regarding the Category 1 problems described in Section 11.1.

11.4.1 <u>Decision Rule</u>

The following rule applies to decisions regarding Problem 1:

If all measured concentrations in all surface and subsurface soil samples collected from a

targeted area are less than background concentrations (if any, see Section 11.4.2) and less than

the screening levels below, then the risk evaluation is complete and there is no unacceptable risk

from the area. In this case, present the data and the risk evaluation in a Data Gaps Assessment

(DGA) report.

It must be noted that screening levels for the decision rules described in this section are different

than PSLs described in Section 11.2.3 (which are used to set desired laboratory quantitation

limits). Soil screening levels are defined as the lowest of the applicable human health risk-

screening criteria, and the selected ecological soil screening levels (SSLs), (for surface soil only).

Furthermore, because the toxicity of individual dioxin and furan congeners vary widely, the

associated screening levels will be the PSLs for total toxicity equivalency (TEQ) of 2,3,7,8-TCDD.

The measured concentrations to be compared with the PSLs will be the total TEQs, calculated by

multiplying each dioxin and furan congener concentration by that congener's toxicity equivalency

factor (TEF) and summing the results. The total TEQ PSL is the same as the PSL presented in

Worksheet #15 for 2,3,7,8-TCDD, which has a TEF of 1. (The individual congener PSLs

presented in Worksheet #15 will not be compared with individual congener concentrations to

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make project decisions in accordance with this decision rule. These individual PSLs are presented to provide approximate values for the evaluation of analytical sensitivity only). If

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concentrations of surface and subsurface soil samples collected from a targeted area are greater

than surface and subsurface soil screening levels, perform the risk screening evaluation. The

SASE screening evaluation will be prepared with figures and tables, including the presentation of

all exceedances of the applicable human health and ecological criteria identified for surface and

subsurface soil (screening levels), and a meeting will be convened with the Project Team to

discuss the next steps to be taken. Note: The screening levels are the same as the PSLs

(Section 11.2.3) with the exception that the screening levels do not include the protection of

groundwater PSLs and the RIDEM criterion.

Note: If exceedances are "serious enough," the project team will tend to recommend a baseline

human health and ecological risk assessment to quantify possible unacceptable levels of

exposure to contaminants. In this instance, the risk assessment will be performed and included

in the DGI report.

11.4.2 <u>Background Comparisons</u>

Comparisons to background soil concentrations will be used to evaluate metals contamination. Metals

commonly occur due to their presence in soil, attributable to geologic conditions.

The background dataset for metals is in the Basewide Background Study Report for Naval Station

Newport (Background Study) (Tetra Tech, July 2008). The method used for comparison between

datasets for metals is outlined in the Background Study. For metals, when the soil type present at the site

can be determined or matched to a particular soil type considered in the background study, a standard

comparison can be made using 95% upper concentration limit (UCL) of the two data sets.

11.5 SPECIFY PERFORMANCE CRITERIA

The sample locations were selected based on the need to characterize the nature and extent of

contamination at the Site. The soil and groundwater analytical data will be used to map the spatial

boundaries of soil and groundwater containing contaminant concentrations exceeding PSLs. Particular

scrutiny will be applied to analytical results below the LOQ when PSLs are below the LOQ. The data

usability evaluation process is described in more detail in Worksheet #37.

The data collected under this SAP are anticipated to be sufficient to delineate the nature and extent of

contamination and support potential baseline risk assessments for the Site. The project team will review

the data as part of the data usability assessment described in Worksheet #37. If any significant data

gaps are identified, the Project Team will determine the next appropriate step.

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### 11.6 **DATA COLLECTION PLAN**

The plan for data collection is provided in detail in Worksheet #17.

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## SAP Worksheet #12 -- Measurement Performance Criteria Table (note matrix in table entry)

(UFP-QAPP Manual Section 2.6.2)

## Measurement Performance Criteria Table – Field QC Samples<sup>(1)</sup>

QC Sample	Analytical Group	Frequency	Data Quality Indicators (DQIs)	Measurement Performance Criteria	QC Sample Assesses Error for Sampling (S), Analytical (A) or both (S&A)
Equipment Rinsate Blank <sup>(2)</sup>	All analytical groups	One per 20 samples	Accuracy /Bias/ Contamination	No target analytes > ½ LOQ (>LOQ for common laboratory contaminants), unless target analytes in field samples are > 10x those in rinsate blank.	S & A
Field Duplicates	Organics	One per 10 samples	Precision	Soils: Relative percent difference (RPD) must be ≤ 50%. Waters: RPD must be ≤ 30%.  If sample results are < 2x LOQ, professional judgment is used.	S & A
	Metals	One per 10 samples	Precision	For values ≥ 5x LOQ Soils: RPD must be ≤ 50% Waters: RPD must be ≤ 30%.  For values < 5x LOQ Soils: Absolute difference must be ≤ 4x LOQ Waters: Absolute difference must be ≤ 2x LOQ for waters.	S & A
Temperature Blank	All analytical groups <sup>(3)</sup>	One per cooler	Representativeness	Temperature must be ≤ 6 °C	S

The measurement performance criteria (MPCs) for laboratory QC samples are presented in Worksheet #28.
 Equipment rinsate blanks will be collected if non-dedicated sampling equipment is used. 3. For metals, the properties of the collected in the collecte For metals, the MPC is only applicable for mercury.

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## SAP Worksheet #13 -- Secondary Data Criteria and Limitations Table

(UFP-QAPP Manual Section 2.7)

Secondary Data	Data Source	Data Generator(s)	How Data Will Be Used	Limitations on Data Use
Report	Site Investigation Report Tank Farm No.2. May 1998	GZA GeoEnvironmental, Inc.	Boring logs will be used when interpreting geologic and hydrogeologic data for the site. Some soil data, and groundwater data will be used for determinations of nature and extent of contamination.	No limitations are applicable.
Report	Draft Work Plan for Site Closure, Tank Farm 2. September 2003.	Foster Wheeler Environmental Corporation.	Some soil and groundwater data will be used for determinations of nature and extent of contamination.	No limitations are applicable.
Report	Draft Site Investigation and Remedial Action Report (SIRAR) for Tank Farm 2. July 2006.	Tetra Tech EC, Inc.	Some soil and groundwater data will be used for determinations of nature and extent. AOC locations and history are being used to determine sample locations, size and analytes.	No limitations are applicable.
Report	Addendum 1, Site Investigation and Remedial Action Report for Tank Farm 2. April 2009.	Tetra Tech EC, Inc.	Some soil data will be used for determinations of nature and extent.	No limitations are applicable.
Report	Groundwater Monitoring Report for Tank Farm 2. August 2009.	Tetra Tech EC, Inc.	Some LNAPL and groundwater data will be used for determinations of nature and extent.	No limitations are applicable.
Report	Basewide Background Investigation Report, Naval Station Newport, Newport Rhode Island. July 2008.	Tetra Tech	Data will be used to determine if metals present onsite are naturally occurring or a result of historic site activities.	No limitations are applicable.

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SAP Worksheet #14 -- Summary of Project Tasks

(UFP-QAPP Manual Section 2.8.1)

The following project tasks are summarized in the sections below:

Field Tasks

Analytical Tasks

Data Management and Review

Project Report

The Tetra Tech and USEPA standard operating procedures (SOPs) and field documentation forms

referred to in this worksheet are included in Appendix C and Appendix D, respectively. Project-specific

procedures for select field tasks are also provided in Appendix E. The field team will follow the project-

specific field procedures unless these procedures do not provide guidance on a specific field task issue.

In that case, the procedures in the cited SOPs will be followed.

14.1 FIELD TASKS

This project will include the following field tasks:

• Mobilization/Demobilization and Utility Clearance - includes mobilization of equipment and staff to the

site, field team orientation, a site walkover, utility clearance, and demobilization. A DIGSAFE number

and NTEC Newport utility clearance will be obtained prior to mobilizing drilling equipment. Detailed

procedures for mobilization are provided in Appendix E-1.

<u>Drilling and Soil Sample Collection</u> – Soil borings will be advanced for continuous soil sampling using

drilling methods described in SOP GH-1.3, or direct-push technology described in SOP SA-2.5.

Boring logs will be created according to SOP GH-1.5. Surface and subsurface soil samples for

laboratory analysis will be collected from the borings according to SOP SA-1.3. Project-specific

procedures for drilling and soil sampling are presented in Appendix E-2. The soil samples will be

collected from the vadose zone at the intervals listed in Worksheet #18.

Field Quality Control Samples – Field quality control (QC) samples will be collected as part of the

investigation, including field duplicates, trip blanks, and equipment rinsate blanks. Worksheet #20

presents the field QC sample summary.

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Field samples to be used for laboratory QC analyses will be assigned by the field sampler on the

chain-of-custody form and sample log sheet. The laboratory will perform matrix spike (MS) and

matrix spike duplicate (MSD) analyses for organic analyses and MS and laboratory duplicate

analyses for metals analysis. Additional sample volume will be collected as necessary for the

laboratory QC analyses.

Field Instrument Calibration - These procedures are described in Worksheet #22.

Equipment decontamination - All non-disposable equipment that comes in contact with the sample

medium will be decontaminated according to SOP SA-7.1 to prevent cross-contamination between

sampling points. This includes equipment such as stainless steel bowls, scoops, as well as heavy

equipment. Personnel decontamination is discussed in the HASP.

All heavy equipment, including the drilling rig, rods and augers, and other down-hole equipment used

during site investigation activities, will be decontaminated prior to beginning work and between all

boreholes using a high-pressure steam wash. Potable water will be used for steam-cleaning.

Investigation-Derived Waste (IDW) Characterization and Disposal – IDW includes decontamination

fluid, used personal protective equipment (PPE), used sampling equipment, and drill cuttings and

excess soil samples. IDW characterization and disposal will be performed after all IDW has been

containerized. IDW shall be handled in accordance with SOP SA-7.1. A summary is presented in

Appendix E-3.

Land Surveying - After completion of sample collection, the coordinates of all sample points,

including soil borings, monitoring wells, as well as other pertinent features will be determined by a

Rhode Island registered land surveyor. The coordinates of the features will be incorporated into the

NAVSTA Newport geographic information system (GIS) database and used for site mapping. Details

of land surveying are presented in Appendix E-4.

14.2 **ANALYTICAL TASKS** 

Chemical analysis of soil samples will be performed by subcontracted laboratories. Test America will

perform the dioxin analysis and Katahdin will perform PAH, PCB, TPH (as GRO and ExTPH) and metals

analyses in accordance with the methods identified in Worksheet #19 and the requirements of the

analytical specifications for laboratory services developed for this work by Tetra Tech (Appendix F).

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TestAmerica and Katahdin will follow the laboratory-specific SOPs (Worksheets #19 and #23) developed, based on the methods listed in Worksheet #19. Copies of the Laboratory SOPs are included in Appendix

G.

All soil sample analytical results will be reported by the laboratory on a dry-weight basis. Results of

percent moisture will be reported in each analytical data package and electronic data files. This

information will also be captured in the project database, which will eventually be uploaded to NIRIS.

Percent moisture information will also be captured in the Data Gaps Reports.

The analytical data packages provided by Test America and Katahdin will be in a contract laboratory

program-like format and will be fully validatable and contain raw data, summary forms for all sample and

laboratory method blank data, and summary forms containing all method specific quality control (results,

recoveries, relative percent differences, relative standard deviations, and/or percent differences etc.).

14.3 DATA MANAGEMENT

Data management will be performed in accordance with SOP CT-05. Data management procedures will

include the following:

• Project documentation and records

Field sample collection and field measurement records are described in Worksheets #27 and

#29.

Laboratory data package deliverables are described in the analytical specifications.

Data assessment documents and records are listed in Worksheet #29.

<u>Data recording formats</u> are described in Worksheet #27.

<u>Data handling and management</u> - After the field investigation is completed, the field sampling log

sheets will be organized by date and media and filed in the project files. The field logbooks for this

project will be used only for this Site, and will also be categorized and maintained in the project files

after the completion of the field program. Project personnel completing concurrent field activities may

maintain multiple field logbooks. When possible, logbooks will be segregated by sampling activity.

The field logbooks will be titled based on date and activity. The data handling procedures to be

followed by the laboratories will meet the requirements of the technical specification. The electronic

data results will be automatically downloaded into the Tetra Tech database in accordance with

proprietary Tetra Tech processes.

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<u>Data tracking and control</u> - The Tetra Tech PM (or designee) is responsible for the overall tracking

and control of data generated for the project.

o Data Tracking. Data is tracked from its generation to its archiving in the Tetra Tech project-

specific files. The Project Chemist (or designee) is responsible for tracking the samples collected

and shipped to the contracted laboratory. Upon receipt of the data packages from the analytical

laboratory, the Project Chemist will oversee the data validation effort, which includes verifying that

the data packages are complete and that results for all samples have been delivered by the

analytical laboratory.

Data Storage, Archiving, and Retrieval. The data packages received from the subcontract

laboratory are tracked in the data validation log book. After the data are validated, the data

packages are entered into the Tetra Tech CLEAN file system and archived in secure files. The

field records including field logbooks, sample logs, chain-of-custody records, and field calibration

logs will be submitted by the FOL to be entered into the CLEAN file system prior to archiving in

secure project files. The project files are audited for accuracy and completeness. At the

completion of the Navy contract, the records will be stored by Tetra Tech.

Data Security. The Tetra Tech project files are restricted to designated personnel only. Records

can only be borrowed temporarily from the project file using a sign-out system. The Tetra Tech Data

Manager maintains the electronic data files. Access to the data files is restricted to qualified

personnel only. File and data backup procedures are routinely performed.

14.4 DATA REVIEW

Data review is described in other worksheets, as follows:

Data verification is described in Worksheet #34.

• Data validation is described in Worksheets #35 and #36.

Usability assessment is described in Worksheet #37.

14.5 PROJECT REPORT

Following completion of the investigations outlined in this SAP, the Navy will prepare a Draft SASE

Report, or a Draft RI Report, in accordance with the decision rules in Worksheet #11.4. This document

will summarize the investigation activities; describe any issues encountered in the field and corrective

actions taken; provide tables comparing soil and groundwater sampling results to screening criteria,

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defined in Section 11.4.1; and provide figures depicting the locations sampled and the spatial distribution

of contaminants. The report will also provide the appropriate risk analysis as described in Worksheet

11.4. The Draft Technical Report will also contain recommendations for the next steps for the Site based

on these analyses.

The Draft Technical Report will be submitted to RIDEM and the USEPA for review. Upon receipt of

regulatory comments, a response will be prepared, and if warranted, a meeting or conference call will be

held to resolve comments. A Final Technical Report incorporating comments will be issued for inclusion

in the NAVSTA Newport Administrative Record.

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

## SAP Worksheet #15 - Reference Limits and Evaluation Table

(UFP-QAPP Manual Section 2.8.1)

In Worksheet #15, the Project Screening Level (PSL) is presented in bold font if it is less than the LOQ but greater than or equal to the LOD; and the PSL is presented as bolded and shaded if it is less than the LOD. The limitations on data usability due to unmet sensitivity goals will be evaluated as described in Worksheet #37 and discussed in the project report.

Matrix: Soil

		Anglytical	Soil PSL <sup>(3)</sup>	Soil	LOQ	Ka	atahdin Lir	nits <sup>(5)</sup>
Analyte <sup>(1)</sup>	CAS Number	Analytical Method <sup>(2)</sup>	(mg/kg)	PSL Reference <sup>(4)</sup>	Goal (mg/kg)	LOQ (mg/kg)	LOD (mg/kg)	DL (mg/kg)
Polycyclic Aromatic Hydrocarbons								
2-Methylnaphthalene	91-57-6	8270D SIM	29	Eco SSL	9.7	0.02	0.01	0.0022
Acenaphthene	83-32-9	8270D SIM	20	Eco SSL	6.7	0.02	0.01	0.0015
Acenaphthylene	208-96-8	8270D SIM	23	Res DEC	7.7	0.02	0.01	0.0012
Anthracene	120-12-7	8270D SIM	29	Eco SSL	9.7	0.02	0.01	0.0012
Benzo(a)anthracene	56-55-3	8270D SIM	0.15	RSL Res	0.05	0.02	0.01	0.0019
Benzo(a)pyrene	50-32-8	8270D SIM	0.015	RSL Res	0.005	0.02	0.01	0.0033
Benzo(b)fluoranthene	205-99-2	8270D SIM	0.15	RSL Res	0.05	0.02	0.01	0.0024
Benzo(g,h,i)perylene	191-24-2	8270D SIM	0.8	Res DEC	0.27	0.02	0.01	0.002
Benzo(k)fluoranthene	207-08-9	8270D SIM	0.9	Res DEC	0.3	0.02	0.01	0.0031
Chrysene	218-01-9	8270D SIM	0.4	Res DEC	0.13	0.02	0.01	0.0017
Dibenzo(a,h)anthracene	53-70-3	8270D SIM	0.015	RSL Res	0.005	0.02	0.01	0.0018
Fluoranthene	206-44-0	8270D SIM	20	Res DEC	6.7	0.02	0.01	0.0018
Fluorene	86-73-7	8270D SIM	28	Res DEC	9.3	0.02	0.01	0.0032
Indeno(1,2,3-c,d)pyrene	193-39-5	8270D SIM	0.15	RSL Res	0.05	0.02	0.01	0.0019
Naphthalene	91-20-3	8270D SIM	3.6	RSL Res	1.2	0.02	0.01	0.0026
Phenanthrene	85-01-8	8270D SIM	29	Eco SSL	9.7	0.02	0.01	0.0018
Pyrene	129-00-0	8270D SIM	1.1	Eco SSL	0.37	0.02	0.01	0.0021
Metals								
Aluminum	7429-90-5	6020A	50	Eco SSL	17	30	4	0.51
Antimony	7440-36-0	6020A	0.27	Eco SSL	0.09	0.1	0.05	0.020
Arsenic	7440-38-2	6020A	0.39	RSL Res	0.13	0.5	0.4	0.15
Barium	7440-39-3	6020A	330	Eco SSL	110	0.2	0.1	0.037
Beryllium	7440-41-7	6020A	1.5	RIDEM Res DEC	0.5	0.1	0.02	0.0041
Cadmium	7440-43-9	6020A	0.36	Eco SSL	0.12	0.1	0.02	0.0076

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Matrix: Soil

		Analytical	Co:1 DC1 (3)	Soil	LOQ	Katahdin Limits <sup>(5)</sup>		
Analyte <sup>(1)</sup>	CAS Number	Analytical Method <sup>(2)</sup>	Soil PSL <sup>(3)</sup> (mg/kg)	PSL	Goal	LOQ	LOD	DL (mg/kg)
			(ilig/kg)	Reference <sup>(4)</sup>	(mg/kg)	(mg/kg)	(mg/kg)	
Calcium	7440-70-2	6020A				10	8	3.8
Chromium	7440-47-3	6020A	0.29	RSL Res	0.097	0.3	0.2	0.049
Cobalt	7440-48-4	6020A	2.3	RSL Res	0.77	0.1	0.03	0.0054
Copper	7440-50-8	6020A	28	Eco SSL	9.3	0.3	0.2	0.071
Iron	7439-89-6	6020A	200	Eco SSL	66.7	10	6	2.40
Lead	7439-92-1	6020A	11	Eco SSL	3.7	0.1	0.05	0.070
Magnesium	7439-95-4	6020A			-	10	8	1.37
Manganese	7439-96-5	6020A	180	RSL Res	60	0.2	0.1	0.042
Mercury	7439-97-6	7471B	0.1	Eco SSL	0.033	0.033	0.017	0.0052
Nickel	7440-02-0	6020A	38	Eco SSL	13	0.2	0.12	0.026
Potassium	7440-09-7	6020A				100	40	4.6
Selenium	7782-49-2	6020A	0.52	Eco SSL	0.17	0.5	0.3	0.039
Silver	7440-22-4	6020A	4.2	Eco SSL	1.4	0.1	0.04	0.0064
Sodium	7440-23-5	6020A				100	40	2.6
Thallium	7440-28-0	6020A	0.0569	Eco SSL	0.019	0.1	0.04	0.0094
Vanadium	7440-62-2	6020A	2	Eco SSL	0.67	0.5	0.4	0.11
Zinc	7440-66-6	6020A	46	Eco SSL	15	1	0.8	0.13
Petroleum Hydrocarbons								
GRO (C5-C12)		8015C				2.5	2	2
ExTPH (C8-C44)		8015C/FL A Pro Mod				5	3	2.6
TPH			500	Res DEC	170	7.5	5	2
PCBs								
Aroclor 1016	12674-11-2	8082A	0.39	Res RSL	0.13	0.017	0.0085	0.0060
Aroclor 1221	11104-28-2	8082A	0.14	Res RSL	0.047	0.017	0.0085	0.0079
Aroclor 1232	11141-16-5	8082A	0.14	Res RSL	0.047	0.017	0.0085	0.0093
Aroclor 1242	53469-21-9	8082A	0.22	Res RSL	0.073	0.017	0.0085	0.0058
Aroclor 1248	12672-29-6	8082A	0.22	Res RSL	0.073	0.017	0.0085	0.0061
Aroclor 1254	11097-69-1	8082A	0.11	Res RSL	0.037	0.017	0.0085	0.0047
Aroclor 1260	11096-82-5	8082A	0.22	Res RSL	0.073	0.017	0.0085	0.0060

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

Matrix: Soil

	CAC	Analytical	Coil DCI (3)	Soil	1.00.000	Tes	tAmerica Li	
Analyte <sup>(1)</sup>	CAS Number	Analytical Method <sup>(2)</sup>	Soil PSL <sup>(3)</sup> (pg/g)	PSL Reference <sup>(4)</sup>	LOQ Goal (pg/g)	LOQ (pg/g)	LOD (pg/g)	MDLs <sup>(6)</sup> (pg/g)
<b>Dioxins and Furans</b>								
1,2,3,4,6,7,8,9-OCDD <sup>(7)</sup>	3268-87-9	8290	14000	So/Air SSL	4700	10	1.5	EDL
1,2,3,4,6,7,8,9-OCDF <sup>(7)</sup>	39001-02-0	8290	14000	So/Air SSL	4700	10	1.5	EDL
1,2,3,4,6,7,8-HPCDD <sup>(7)</sup>	35822-46-9	8290	420	So/Air SSL	140	5	0.75	EDL
1,2,3,4,6,7,8-HPCDF <sup>(7)</sup>	67562-39-4	8290	420	So/Air SSL	140	5	0.75	EDL
1,2,3,4,7,8,9-HPCDF <sup>(7)</sup>	55673-89-7	8290	420	So/Air SSL	140	5	0.75	EDL
1,2,3,4,7,8-HXCDD <sup>(7)</sup>	39227-28-6	8290	42	So/Air SSL	14	5	0.75	EDL
1,2,3,4,7,8-HXCDF <sup>(7)</sup>	70648-26-9	8290	42	So/Air SSL	14	5	0.75	EDL
1,2,3,6,7,8-HXCDD <sup>(7)</sup>	57653-85-7	8290	42	So/Air SSL	14	5	0.75	EDL
1,2,3,6,7,8-HXCDF <sup>(7)</sup>	57117-44-9	8290	42	So/Air SSL	14	5	0.75	EDL
1,2,3,7,8,9-HXCDD <sup>(7)</sup>	19408-74-3	8290	42	So/Air SSL	14	5	0.75	EDL
1,2,3,7,8,9-HXCDF <sup>(7)</sup>	72918-21-9	8290	42	So/Air SSL	14	5	0.75	EDL
1,2,3,7,8-PECDD <sup>(7)</sup>	40321-76-4	8290	4.2	So/Air SSL	1.4	5	0.75	EDL
1,2,3,7,8-PECDF <sup>(7)</sup>	57117-41-6	8290	140	So/Air SSL	46.7	5	0.75	EDL
2,3,4,6,7,8-HXCDF <sup>(7)</sup>	60851-34-5	8290	42	So/Air SSL	14	5	0.75	EDL
2,3,4,7,8-PECDF <sup>(7)</sup>	57117-31-4	8290	14	So/Air SSL	4.7	5	0.75	EDL
2,3,7,8-TCDD <sup>(7)</sup>	1746-01-6	8290	4.2	So/Air SSL	1.4	1	0.15	EDL
2,3,7,8-TCDF <sup>(7)</sup>	51207-31-9	8290	42	So/Air SSL	14	1	0.15	EDL
TOTAL HPCDD	37871-00-4	8290						
TOTAL HPCDF	38998-75-3	8290						
TOTAL HXCDD	34465-46-8	8290						
TOTAL HXCDF	55684-94-1	8290						
TOTAL PECDD	36088-22-9	8290						
TOTAL PECDF	30402-15-4	8290						
TOTAL TCDD	41903-57-5	8290						
TOTAL TCDF	55722-27-5	8290						

Site Name: Tank Farm 2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

### Notes for Worksheet 15:

- 1. Soil samples will be analyzed for PAHs, metals, dioxins (select samples) and PCBs (select samples).
- All methods are EPA SW-846.
- 3. Although there are separate PSLs for surface and subsurface soil, a single soil PSL representing the lowest of these PSLs is presented here, and the LOQ goals and selected methods are the same for all soil samples, in order to simplify sampling and analysis procedures. The soil PSLs presented are the lowest of:
  - EPA Regional Screening Levels (RSLs) residential and industrial soil values (EPA, 2010a)
  - EPA Soil to Air Soil Screening Levels (SSLs) (EPA, 2010b)
  - RIDEM Residential Direct Exposure Criteria (RIDEM, 2011)
  - Selected ecological SSL (applicable only for surface soil PSLs)

One-tenth values are displayed for non-cancer RSLs and Soil to Air SSLs to correspond to a target hazard quotient of 0.1. The selected ecological SSLs are the lowest of the selected benchmarks for plants, invertebrates, and wildlife. The benchmarks were selected by order of preference according to the following hierarchy:

Order of preference for plants and invertebrates:

- 1. EPA Ecological SSLs (U.S. EPA, 2003-2008)
- 2a. Oak Ridge National Laboratory (ORNL) Plant Toxicological Benchmark (Efroymson, 1997a)
- 2b. ORNL Invertebrate Toxicological Benchmark (Efroymson, 1997b)
- 3. Canadian Council and Ministers of Environment (CCME) (CCME, 1997-2010)
- 4. Target values for soil remediation (MHSPE, 2000)

Order of preference for wildlife:

- 1. EPA Ecological SSLs (U.S. EPA, 2003-2008)
- 2. CCME (CCME, 2010)
- 4. EPA Region 5 Ecological Screening Levels (U.S. EPA, 2003).4.

**PSL** Reference Abbreviations:

- Eco SSL = Selected ecological SSL
- Res RSL = EPA RSL residential soil value
- So/Air SSL = EPA Soil to Air SSL
- Res DEC = RIDEM Residential Direct Exposure Criteria
- 5. The LOQs, LODs, and DLs presented for metals analyzed by Method 6020A reflect an assumed dilution factor of 5, which is typically required for solids analysis by this method. If the dilution factor is different, the values will be adjusted accordingly.
- 6. Estimated Detection Limit (EDL) For each chemical not detected, an EDL is calculated. The sample-specific EDL is an estimate made by the laboratory of the concentration of a given chemical that would have to be present to produce a signal with a peak height of at least 2.5 times the background signal level. The estimate is specific to a particular analysis of the sample and will be affected by sample size, dilution, and so forth. Non-detected results will be reported with an associated value of the EDL, and results between the LOQ and EDL will be flagged as estimated "J". LODs are presented for informational purposes.
- 7. PSL value presented is the screening level for total toxicity equivalency (TEQ) of 2,3,7,8-TCDD (4.2 pg/g), divided by the congener's 2005 World Health Organization (WHO) toxicity equivalency factor (TEF) for humans and mammals (Van den Berg, et al, 2006). This value is presented as an approximate value by which to evaluate analytical sensitivity, but it will not be compared with the individual dioxin or furan congener's concentrations to make project decisions according to the decision rules described in Worksheet #11, Section 11.4. To make the project decisions, each congener concentration will be multiplied by the congener's TEF; the TEF-adjusted concentrations of all congeners will be summed to obtain the total TEQ, and the total TEQ will be compared with the total TEQ PSL.

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# SAP Worksheet #16 -- Project Schedule / Timeline Table (optional format) (UFP-QAPP Manual Section 2.8.2)

		Dates (M	M/DD/YY)		Deliverable	
		Anticipated Date(s) of Initiation	Anticipated Date of Completion	Deliverable	Deliverable Due Date	
Soil sampling	Tetra Tech	July , 2013	August, 2013	Draft Data Gaps Assessment Report	January, 2014	

Site Name: Tank Farms 2, NAVSTA Newport Project Name: Data Gaps Investigation

Site Location: Portsmouth, Rhode Island

Title: Sampling and Analysis Plan Document No.: W5211722F Revision Number: 0

Date: July 2013

SAP Worksheet #17 -- Sampling Design and Rationale

(UFP-QAPP Manual Section 3.1.1)

The sampling design for this project is based on the need to fill data gaps that exist after the completion

of various other site investigations and remedial actions by the DESC. To fill the data gaps Tetra Tech

has expanded the list of analytes and the sampling areas in places to be investigated under this SAP.

Tank Sludge Burning AOCs: Areas that were AOCs in a former site investigation (TtEC, 2006) (AOC-

001; -003; -004 and -005) and were originally screened and tested for a limited number of analytes in soil,

mainly TPH using Petroflag™ screening and/or laboratory analyses. These areas were reported to have

open-burning of tank sludge, as determined from aerial photographs taken during the 1950's. Given the

site history, it has been decided to expand the analyte list to include contaminants that may be present

due to burning sludge and expand the sampling area and the number of soil samples being collected in

these areas. Groundwater samples will not be collected in these areas because groundwater has been

monitored (historically at wells GZ-201, GZ-203, GZ-225, GZ-226, GZ-227 and RW-2) and results did not

suggest contamination migration from soil to groundwater.

The soil sampling areas are designed to be approximately 50 percent larger than the AOC that was

defined, based on aerial photography review (TtEC, 2006). The reason is to increase the chance that the

extent of contamination that may be present is determined. The sampling area for each AOC is small and

contains a larger area of potentially contaminated soil, thereby making the sampling area conservative.

Sample locations will be determined using a 15 by 15 foot grid superimposed over the sampling area.

This strategy increases the density and spatial distribution of sampling within the AOC compared to the

previous investigation (TtEC, 2006). Additional sampling locations were added in the center of each grid

square at AOC-001 and -003 to increase the number of samples located within the AOC boundary.

Sampling locations will be located in the field using a global positioning system (GPS). The extent of the

old sampling area, the new sampling area, and the location of samples that will be collected are shown on

Figures 6, 7 and 8.

Soil borings will be advanced using direct push technology (DPT) or conventional drilling methods. Soil

samples will be collected from the 0 to 1, 2 to 4, and 8 to 10 foot intervals at AOC-001;-003; -004 and -

005 assuming bedrock is deeper than 10 feet. If bedrock is encountered prior to reaching a depth of 10

feet, it will be at the sampler's discretion, based on depth of refusal, as to how many analytical samples

are collected at the soil boring. Based on previous investigations, shallow bedrock (less than five feet)

was encountered across the AOC-004 and -005 areas and it is therefore expected that two samples will

therefore be collected from borings in these AOCs. However, at the request of the EPA, allowance for

three samples remain herein in case bedrock is deeper. A sample from directly above bedrock will be

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Site Name: Tank Farms 2, NAVSTA Newport Project Name: Data Gaps Investigation

Site Location: Portsmouth, Rhode Island

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collected from each location. Soil samples will be analyzed for PAHs, metals, and dioxins from the 0 to 4

foot range and PAHs, and metals below 4 feet. At the request of the RIDEM, up to ten percent of the soil

samples collected for analysis from the sludge burning AOCs will be analyzed for TPH (GRO and

ExTPH). Samples will be submitted for GRO and ExTPH analyses if there is visual and/or olfactory

evidence of petroleum contamination observed in the field. The number of sampling locations has been

summarized in Table 17-1. The locations where field duplicate and QC samples are collected will be

determined in the field by the FOL.

**Building 219** 

Soil samples will be collected for PCB analysis using DPT drilling methods at Building 219 (Figure 5).

Surface soil samples will be collected from 0 to 1 feet bgs, and subsurface samples will be collected from

2-4 feet bgs. Sixteen samples (eight surface soil and eight subsurface soil, at the depths described

above) will be collected at the four locations that were analyzed in 2005, See Figure 5 for details.

**Former Buoy Storage Area** 

Soil samples will be collected for lead analysis using DPT drilling methods at the former buoy storage

area. Surface soil samples will be collected from 0-1 feet bgs, and subsurface samples will be collected

from 2-4 feet bgs. Sixteen samples (eight surface soil and eight subsurface soil) will be collected at eight

locations laid out on a 40 feet by 40 feet grid to cover the areas of concern (Figure 9).

The sampling SOPs for soil sampling are identified in Worksheet #18 and included in Appendix A.

Project-specific sampling procedures are detailed in Appendix C.

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Table 17-1 – Number of Sample Locations by Analytical Group, Matrix, Tank Farm Area, and Depth (Soils)

Category/Area Depth (Soils)	PAHs	Metals	Dioxins	GRO	ExTPH	PCBs	Lead
Soil							
AOC TF2-001							
0-1' bgs	11	11	11	1	1		
2-4' bgs	11	11	11	1	1		
8-10' bgs*	11	11		1	1		
AOC TF2-003		•					
0-1' bgs	9	9	9	1	1		
2-4' bgs	9	9	9	1	1		
8-10' bgs*	9	9		1	1		
AOC TF2-004 and AOC TF2-0	)05	•					
0-1' bgs	49	49	49	5	5		
2-4' bgs	49	49	49	5	5		
8-10' bgs*	49	49		5	5		
Building 219		•					
0-1' bgs						8	
2-4' bgs						8	
Buoy Storage Area							_
0-1' bgs							8
2-4' bgs							8
Total Soils	207	207	138	21	21	16	16

<sup>\*</sup>The sample depth of 8 to 10-ft is based upon a depth to bedrock of 10-feet bgs. One of the samples from each location will be collected from directly above bedrock.

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

# SAP Worksheet #18 -- Sampling Locations and Methods/SOP Requirements Table (UFP-QAPP Manual Section 3.1.1)

Category	Area	Location Identification <sup>(1)</sup>	Matrix	<b>Depth</b> (bgs)	Analytical Group	Number of Samples <sup>(2)</sup>	Sampling SOP Reference <sup>(3)</sup>
				0-1 ft, 2-4 ft	PAHs, Metals, Dioxins	22	
	AOC TF2-001	TF2-SB1000 -		0-1 ft, 2-4 ft	GRO, ExTPH	2	
	AOC 1F2-001	TF2-SB1010		8-10 ft	PAHs, Metals	11	GH-1.3, SA-1.3
1			Soil	8-10 ft	GRO, ExTPH	1	
'		TF2-SB1011 – TF2-SB1019		0-1 ft, 2-4 ft	PAHs, Metals, Dioxins	18	
	AOC TF2-003			0-1 ft, 2-4 ft	GRO, ExTPH	2	
	AOC 1F2-003			8-10 ft	PAHs, Metals	9	
				8-10 ft	GRO, ExTPH	1	

Category	Area	Location Identification <sup>(1)</sup>	Matrix	<b>Depth</b> (bgs)	Analytical Group	Number of Samples <sup>(2)</sup>	Sampling SOP Reference <sup>(3)</sup>
	AOC TF2-004 and AOC TF2-005	and TF2-SB1020 -		0-1 ft, 2-4 ft PAHs, Metals, Dioxins	98		
				0-1 ft, 2-4 ft	GRO, ExTPH	10	
			TF2-SB1068		8-10 ft <sup>(4)</sup>	PAHs, Metals	49
			8-10 ft <sup>(4)</sup>	8-10 ft <sup>(4)</sup>	GRO, ExTPH	5	

Category	Area	Location Identification <sup>(1)</sup>	Matrix	<b>Depth</b> (bgs)	Analytical Group	Number of Samples <sup>(2)</sup>	Sampling SOP Reference <sup>(3)</sup>
1	Building 219	TF2-B219-SB-1080 - TF2-B219-SB-1087	Soil	0-1 ft	PCBs	8	GH-1.3, SA-1.3
1	Building 219	TF2-B219-SB-1080 - TF2-B219-SB-1087	Soil	2-4 ft	PCBs	8	GH-1.3, SA-1.3
1	Buoy Storage Area	TF2-BSA-SB-SS-1090 – TF2-BSA-SB-SS-1097	Soil	0-1 ft	Lead	8	GH-1.3, SA-1.3
1	Buoy Storage Area	TF2-BSA-SB-SB-1090 – TF2-BSA-SB-SB-1097	Soil	2-4 ft	Lead	8	GH-1.3, SA-1.3

Site Name: Tank Farm 2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

## SAP Worksheet #18 -- Sampling Locations and Methods/SOP Requirements Table (Continued)

(UFP-QAPP Manual Section 3.1.1)

### Notes:

- 1. Soil sample ID numbers include a suffix of –NNNN to represent interval depth.
- 2. Field duplicates will be selected based on field conditions at the time of the sampling event.
- 3. Refer to Worksheet #21 for complete reference. SOPs are included in Appendix C. Project-specific sampling procedures are provided in Appendix E.
- 4. If bedrock is encountered prior to reaching a depth of 10 feet, it will be at the sampler's discretion, based on depth of refusal, as to how many analytical samples are collected at the soil boring. Based on previous investigations, shallow bedrock (less than five feet) was encountered across the AOC-004 and -005 areas and it is therefore expected that two samples will therefore be collected from these AOCs. A sample directly above bedrock will be collected at each location.

### Abbreviation:

bgs = below ground surface

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

# SAP Worksheet #19 -- Analytical SOP Requirements Table (UFP-QAPP Manual Section 3.1.1)

Matrix	Analytical Group	Analytical and Preparation Method / SOP Reference <sup>(1)</sup>	Containers <sup>(2)</sup> (number, size, and type)	Sample volume <sup>(3)</sup> (units)	Preservation Requirements (chemical, temperature, light protected)	Maximum Holding Time <sup>(4)</sup> (preparation / analysis)
Soil	GRO	SW-846 5035A, 8015C/ CA-316	Two 40-mL VOC vials	5 g	5 ml methanol; cool to ≤ 6 °C	14 days to analysis
	PAHs	SW-846 3540C or 3550C, 8270D SIM/CA-213, CA-512, CA-526		30 g	Cool to ≤ 6 °C	14 days to extraction; 40 days to analysis
	ExTPH	SW-846 3540C or 3550C, 8015C/ CA-315, CA-527, CA- 535	8-oz wide mouth jar	30 g	Cool to ≤ 6 °C	14 days to extraction; 40 days to analysis
Con	PCBs	SW-846 3540C, 3545A or 3550C, 8082A/CA-329, CA- 500, CA-524, CA-537		30 g	Cool to ≤ 6 °C	30 days to extraction; 40 days to analysis <sup>(6)</sup>
	Metals	SW-846 3050B, 6020A, 7471B/ CA-605, CA-611, CA-627	4-oz wide mouth jar	2 g	Cool to ≤ 6 °C	180 days to analysis except for mercury; 28 days to analysis for mercury
	Dioxins	SW-846 8290/ WS-IDP-0005, WS-ID-0005	One 8-oz glass jar with Teflon®-lined lid or stainless steel liner	30 g	Cool to ≤ 6 °C	30 days to extraction; 45 days to analysis
	GRO	SW-846 5030B, 8015C / CA- 316	Two 40-mL VOC vials	40 mL	HCl to pH < 2, no headspace cool to ≤ 6 oC	14 days to analysis
Aqueous (Rinsate Blanks)	PAHs	SW-846 3510C or 3520C, 8270D SIM/ CA-213, CA-502	Two 1-liter (L) amber glass bottles	1000 ml	Cool to ≤ 6 °C	7 days to extraction; 40 days to analysis
Biarnoj	ExTPH	SW-846 3510C or 3520C, 8015C/ CA-315, CA-520	Two 1-L amber glass bottles	1000 mL	Cool to ≤ 6 oC	7 days to extraction; 40 days to analysis

Site Name: Tank Farm 2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

Matrix	Analytical Group	Analytical and Preparation Method / SOP Reference <sup>(1)</sup>	Containers <sup>(2)</sup> (number, size, and type)	Sample volume <sup>(3)</sup> (units)	Preservation Requirements (chemical, temperature, light protected)	Maximum Holding Time <sup>(4)</sup> (preparation / analysis)
	Dioxins	SW-846 8290/ WS-IDP-0005, WS-ID-0005	Two 1-L Amber Glass Bottles	1000 ml	Cool to ≤ 6 °C	30 days to extraction; 45 days to analysis
	PCBs	SW-846 3510C or 3520C, 8082A CA-329, CA-515	Two 1-L Amber Glass Bottles	1000 ml	Cool to ≤ 6 °C	30 days to extraction; 40 days to analysis
	Metals	SW-846 3010A, 6020A, 7470A/ CA-604, CA-615, CA- 627	500 ml polyethylene bottle	100 ml	Nitric acid to pH < 2, cool to ≤ 6 °C	180 days to analysis except for mercury; 28 days to analysis for mercury

- 1. Refer to the Analytical SOP References table (Worksheet #23).
- 2. Laboratories may provide specific containers at their discretion.
- 3. Minimum sample volume or mass requirement.
- 4. Maximum holding time is calculated from the time the sample is collected to the time the sample is extracted, digested, or analyzed.
- 5. SW-846 8082A does not specify a holding time to extraction; 30 days is presented here as a conservative limit. The method recommends a holding time of 40 days from extraction to analysis for extracts stored under refrigeration in the dark; but it also refers to SW-846 Chapter 4, which specified that there is no holding time for PCBs. Additionally, SW-846 8082A states that the holding times listed in the method under the conditions listed may be as long as a year.

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

## SAP Worksheet #20 -- Field Quality Control Sample Summary Table

(UFP-QAPP Manual Section 3.1.1)

Matrix	Analytical Group	No. of Sampling Locations	No. of Field Duplicates <sup>1</sup>	No. of Assigned Laboratory QC Samples <sup>2</sup>	No. of Field Blanks	No. of Equip. Blanks <sup>3</sup>	No. of PT Samples	Total No. of Samples to Lab <sup>4</sup>
	GRO	21	2	1	0	1	0	24
	ExTPH	21	2	1	0	1	0	24
	PAHs	207	21	10	0	10	0	238
Soil	Dioxins	138	14	7	0	7	0	159
	Metals	207	21	10	0	10	0	238
	Lead	16	2	1	0	1	0	20
	PCBs	16	2	1	0	1	0	20

- 1. Collect 1 field duplicate per 10 field samples for each matrix.
- 2. Assign 1 Laboratory QC sample per 20 samples for MS/MSD analysis for organics and MS/laboratory duplicate analysis for metals. For other soil analytes, no extra volume is required.
- 3. Collect 1 rinsate blank per 20 field samples.
- 4. Total number of samples does not include the laboratory QC samples.

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

## SAP Worksheet #21 -- Project Sampling SOP References Table

(UFP-QAPP Manual Section 3.1.2)

Reference Number	Title, Revision Date and / or Number	Originating Organization of Sampling SOP	Equipment Type	Modified for Project Work?	Comments <sup>1</sup>
CT-05	CT-05 - Database Records and Quality Assurance; Revision 2, 2001	Tetra Tech	Not applicable	No	
GH-1.2	GH-1.2 - Evaluation of Existing Monitoring Wells and Water Level Measurement; Revision 2, 2003	Tetra Tech	Electronic water level indicator	No	Also see Appendix E-2 for project-specific procedures to augment SOP.
GH-1.3	GH-1.3 - Soil and Rock Drilling Methods; Revision 1, 1999	Tetra Tech	Drilling rig and accessories	No	
GH-1.5	GH-1.5 - Borehole and Sample Logging; Revision 1, 1999	Tetra Tech	Not applicable	No	
GH-2.5	GH-2.5 – Groundwater Contour Maps and Flow Determination; Revision 1, 2009	Tetra Tech	Not applicable	No	
GH-2.8	GH-2.8 - Groundwater Monitoring Well Installation; Revision 3, 2003	Tetra Tech	Drilling rig, accessories, and well supplies	No	Also see Appendix E-2 for project-specific procedures to augment SOP.
HS-1.0	HS-1.0 - Utility Locating and Excavation Clearance; Revision 2, 2003	Tetra Tech	Remote subsurface sensing, magnetometer, GPR, etc.	No	
SA-1.3	SA-1.3 - Soil Sampling; Revision 9, 2009	Tetra Tech	Sampling supplies	No	Also see Appendix E-2 for project-specific procedures to augment SOP.
SA-2.5	SA-2.5 – Direct Push Technology	Tetra Tech	Geoprobe®; Hydropunch™	No	
SA-6.1	SA-6.1 - Non-Radiological Sample Handling; Revision 3, 2004	Tetra Tech	Not applicable	No	
SA-6.3	SA-6.3 - Field Documentation; Revision 3, 2009	Tetra Tech	Not applicable	No	
SA-7.1	SA-7.1 - Decontamination of Field Equipment; Revision 6, 2009	Tetra Tech	Not applicable	No	Also see Appendix E-3 for project-specific procedures to augment SOP.
GW 001	Low Stress (Low Flow) Purging and Sampling Procedure for the Collection of Groundwater Samples from Monitoring Wells; Revision 3, 2010 / GW 001	EPA Region 1	Submersible pump	Yes <sup>(2)</sup>	Also see Appendix E-2 for project-specific procedures to augment SOP.

<sup>1.</sup> SOPs are included as Appendix C. Appendix E contains project specific field task procedures, including Appendix E-1 Mobilization, Appendix E-2 Project Specific Sampling Procedures, Appendix, E-3 Investigation Derived Waste Management, and Appendix E-4 Surveying.

2. If saturated screen length is greater than 10 feet, the sampling procedures will be modified as described in Appendix E-2.

# **Project-Specific Sampling and Analysis Plan** Site Name: Tank Farm 2, NAVSTA Newport

Site Name: Tank Farm 2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

# SAP Worksheet #22 -- Field Equipment Calibration, Maintenance, Testing, and Inspection Table (UFP-QAPP Manual Section 3.1.2.4)

Field Equipment	Activity <sup>1</sup>	Frequency	Acceptance Criteria	Corrective Action	Resp. Person	SOP Reference	Comments
Photo-Ionization Detector (PID) /Flame Ionization Detector (FID)	Visual Inspection  Calibration/  Verification	Daily  Beginning and end of day	Manufacturer's guidance	Operator correction or Replacement	Field Operations Leader	Manufacturer's instruction manual	Rental field equipment will be used

<sup>&</sup>lt;sup>1</sup>Rental equipment and instruments will be used in the field. The rental firms will be responsible for the proper care, maintenance, and repair of these items, and for tracking and documenting equipment and instrument maintenance and repairs.

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

# SAP Worksheet #23 -- Analytical SOP References Table (UFP-QAPP Manual Section 3.2.1)

Lab SOP Number	Title, Revision Date, and / or Number	Definitive or Screening Data	Matrix and Analytical Group	Instrument	Organization Performing Analysis	Modified for Project Work? (Y/N)
CA-213	Analysis of SVOC By: SW 846 Method 8270 – Modified For Selected Ion Monitoring (SIM), 04/10, Revision 8.	Definitive	Soil and Water/PAHs	GC/MS	Katahdin Analytical	N
CA-329	Analysis Of PCBs As Total Aroclors By Gas Chromatography/Electron Capture Detector (GC/ECD): SW-846 Method 8082, 04/10, Revision 11.	Definitive	Soil and Water/PCBs	GC/Electron Capture Detector (ECD)	Katahdin Analytical	N
CA-500	Preparation Of Sediment/Soil Samples By Sonication Using Method 3550 For Subsequent Pesticides/PCBs Analysis, 08/10, Revision 7.	Definitive	Soil/PCBs	Not applicable (extraction)	Katahdin Analytical	N
CA-502	Preparation Of Aqueous Samples For Extractable Semivolatile Analysis, 10/09, Revision 6.	Definitive	Water/ PAH Extraction	Not applicable (extraction)	Katahdin Analytical	N
CA-512	Preparation Of Sediment/Soil Samples By Sonication Using Method 3550 For Subsequent Extractable Semi-Volatiles Analysis, 08/10, Revision 8.	Definitive	Soil/ PAH Extraction	Not applicable (extraction)	Katahdin Analytical	N
CA-515	Preparation of Aqueous Samples for Pesticides/PCBs Analysis, 08/10, Revision 7.	Definitive	Water/PCBs	Not applicable (extraction)	Katahdin Analytical	N
CA-524	Preparation Of Sediment/Soil Samples By Soxhlet Extraction Using Method 3540 For Pesticide/PCB Analysis, 08/10, Revision 7.	Definitive	Soil/PCBs	Not applicable (extraction)	Katahdin Analytical	Ν
CA-526	Preparation Of Sediment/Soil Samples By Soxhlet Extraction Using Method 3540 For Subsequent Extractable Semivolatile Analysis, 08/10, Revision 7.	Definitive	Soil/ PAHs Extraction	Not applicable (extraction)	Katahdin Analytical	N

Lab SOP Number	Title, Revision Date, and / or Number	Definitive or Screening Data	Matrix and Analytical Group	Instrument	Organization Performing Analysis	Modified for Project Work? (Y/N)
CA-537	Preparation of Sediment/Soil and Tissue Samples by Accelerated Solvent Extraction Using Method 3545 for Subsequent Extractable Pesticide and PCB Analysis, 12/10, Revision 3.	Definitive	Soil/PCBs	Not applicable (extraction)	Katahdin Analytical	N
CA-604	Acid Digestion of Aqueous Samples by EPA Method 3010 for ICP and ICP-MS Analysis of Total or Dissolved Metals, 04/10, Revision 5.	Definitive	Water/Metals Digestion	Not applicable (digestion)	Katahdin Analytical	N
CA-605	Acid Digestion of Solid Samples by USEPA Method 3050 for Metals by ICP- AES and GFAA, 09/10, Revision 5.	Definitive	Soil/Metals Digestion	Not applicable (digestion)	Katahdin Analytical	N
CA-611	Digestion and Analysis of Solid Samples for Mercury by USEPA Method 7471, 12/10, Revision 8.	Definitive	Soil/Mercury	Mercury Analyzer	Katahdin Analytical	N
CA-615	Digestion and Analysis of Aqueous Samples for Mercury by USEPA Method 7470, 04/10, Revision 5.	Definitive	Water/Mercury	Mercury Analyzer	Katahdin Analytical	N
CA-627	Trace Metals Analysis By ICP-MS Using USEPA Method 6020, 04/10, Revision 7.	Definitive	Soil and Water/Metals	ICP-MS	Katahdin Analytical	N
WS-ID-0005	Analysis of Samples for Polychlorinated Dioxins and Furans by HRGC/HRMS 12/09, Revision 7.3.	Definitive	Soil and Water/ Dioxins	GC/HRMS	TestAmerica West Sacramento	N
WS-IDP-0005	Preparation of Samples for Analysis of Polychlorinated Dioxins and Furans for Analysis HRGC/HRMS, 02/10, Revision 1.1.	Definitive	Soil and Water/ Dioxins Extraction	Not applicable (extraction)	TestAmerica West Sacramento	N

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

# SAP Worksheet #24 -- Analytical Instrument Calibration Table (UFP-QAPP Manual Section 3.2.2)

Instrument	Calibration Procedure	Frequency of Calibration	Acceptance Criteria	Corrective Action	Person Responsible for Corrective Action	SOP Reference
GC/MS (SIM) PAHs	Decafluorotriphe nyl-phosphine (DFTPP) Tune. Also DDT, pentachlorophen ol and benzidine check for injection port inertness and GC column performance.  Beginning of each analytical run or every 12 hours  12 hours  13 every 12 hours		DFTPP within method specifications for tuning criteria.  DDT degradation should be < 20%.  Pentachlorophenol and benzidine responses should not exceed a tailing factor of 2 as per Section 11.3.1.3 of Method 8270D.	Re-tune instrument, clean MS source as needed.	Analyst, Department Manager	CA-213
	ICAL - 5-7 (minimum 5 points required) calibration standards, initial calibration for all analytes.	Instrument receipt, major instrument change, when CCV does not meet criteria.	Project specific criteria: Average RF for all PAHs must be $\geq$ 0.05.  Percent Relative Standard Deviation (%RSD) for RFs for all PAHs must be $\leq$ 30% or one option below: Option 1) Linear least squares regression: correlation coefficient (r) must be $\geq$ 0.99 Option 2) Non-linear regression: coefficient of determination ( $r^2$ ) must be $\geq$ 0.99 (6 points for second order).	Recalibrate and/or perform the necessary equipment maintenance. Check the calibration standards. Reanalyze the affected data.	Analyst, Department Manager	
	Initial Calibration Verification (ICV) (Second Source)	Once after each ICAL.	Percent recovery (%R) must be within 80-120% for all project compounds.	Correct problem and verify second source standard. Rerun second source verification. If that fails, correct problem and repeat ICAL.	Analyst, Department Manager	
	Establish RT Window Position	Once per ICAL for each analyte and surrogate.	Position shall be set using the midpoint standard of the ICAL curve when ICAL is performed. On days when ICAL is not performed, the initial CCV is used.	Not applicable.	Analyst, Department Manager	

Instrument	Calibration Procedure	Frequency of Calibration	Acceptance Criteria	Corrective Action	Person Responsible for Corrective Action	SOP Reference
GCMS - PAHs (cont.)	RRTs	With each sample.	RRT of each target compound must be within ±0.06 RRT units.	Correct problem, then rerun ICAL.	Analyst, Department Manager	
	CCV	Daily before sample analysis and every 12 hours	Project specific criteria: For all PAHs RF must be ≥ 0.05  All PAHs and surrogates must be ≤ 25%D  (D = Difference or Drift);	DoD project level approval must be obtained for each of the failed analytes or corrective action must be taken. Correct problem, then rerun calibration verification. If that fails, then repeat ICAL. Reanalyze all samples since last acceptable CCV.	Analyst, Department Manager	
ICP-MS	Tune	Daily prior to calibration	Mass calibration must be within 0.1 amu of true value, Resolution must be < 0.9 amu at 10% peak height.  RSD must be ≤ 5% for at least four replicate analyses.	Perform necessary equipment maintenance.	Analyst, Department Manager	CA-627
	ICAL	Daily prior to sample analysis.	4 point calibration plus blank – The r must be ≥ 0.995.	Recalibrate and/or perform necessary equipment maintenance. Check calibration standards.	Analyst, Department Manager	
	ICV (Second Source)	Once after each ICAL, and before beginning a sample run.	The %R must be within 90-110% for all analytes.	Do not use results for failing elements unless the ICV > 110% and the sample results are non-detect. Investigate and correct problem.	Analyst, Department Manager	
	Calibration Blank	Before beginning a sample sequence, after every 10 samples and at end of the analysis sequence.	No analytes detected > LOD. For negative blanks, absolute value < LOD.	Correct the problem, then reprepare and reanalyze.	Analyst, Department Manager	

Instrument	Calibration Procedure	Frequency of Calibration	Acceptance Criteria	Corrective Action	Person Responsible for Corrective Action	SOP Reference
ICP-MS (cont.)	CCV	After every 10 samples and at the end of each run sequence.	The %R must be within 90-110% for all analytes.	Correct problem, rerun calibration verification. If that fails, then repeat ICAL. Reanalyze all samples since the last successful calibration verification.	Analyst, Department Manager	
	Low-level Calibration Check Standard	ICAL.	The %R must be within 80-120% for all analytes.	Do not use results for failing elements, unless low-level standard recovery.> upper limit and sample results are non-detect. Investigate and correct the problem.	Analyst, Department Manager	
	Interference Check Sample (ICS) - ICSA & ICSB	Daily, before sample injections	The absolute value of the ICSA concentration for all non-spiked analytes (except verified trace impurities) must be less than the LOD (2), and ICSB %Rs must be within 80-120%.	Correct the problem, then reprepare checks and reanalyze all affected samples.	Analyst / Supervisor	
Mercury analyzer	ICAL	Upon instrument receipt, major instrument change, at the start of each day.	Initial Calibration, 5 points plus a calibration blank - $r \ge 0.995$ .	Recalibrate and/or perform necessary equipment maintenance. Check calibration standards.	Analyst, Department Manager	CA-611, CA- 615
	ICV (Second Source)	Once after each ICAL, prior to beginning a sample run.	The %R must be within 90-110% for mercury.	Correct problem and verify second source standard. Rerun ICV. If that fails, correct problem and repeat ICAL.	Analyst, Department Manager	
	Calibration Blank	Before beginning a sample sequence, after every 10 samples and at end of the analysis sequence.	No analytes detected > LOD. For negative blanks, absolute value must be < LOD.	Correct problem. Re-prep and reanalyze calibration blank. All samples following the last acceptable calibration blank must be reanalyzed.	Analyst, Department Manager	
	CCV	Beginning and end of each run sequence and every 10 samples.	The %R must be within 80-120% for mercury.	Correct problem, rerun calibration verification. If that fails, then repeat ICAL. Reanalyze all samples since the last successful calibration verification.	Analyst, Department Manager	

Instrument	Calibration Procedure	Frequency of Calibration	Acceptance Criteria	Corrective Action	Person Responsible for Corrective Action	SOP Reference
GC/HRMS	Tune / Mass Resolution Check (PFK)	At the beginning and the end of each 12- hour period of analysis.	Resolving power ≥ 10,000 at m/z=304.9842 & m/z=380.9760 + 5ppm of expected mass. Lockmass ion between lowest and highest masses for each descriptor and level of reference ≤ 10% full-scale deflection.	Retune instrument & verify. Assess data for impact if end resolution is less than 10,000 narrate or re-inject as necessary.	Analyst /Lab Manager	WS-ID-0005
HRGC/HRMS	GC Column Performance Check (CPSM/WDM per method)	Prior to ICAL or calibration verification.	Peak separation between 2,3,7,8-TCDD and other TCDD isomers result in a valley of ≤ 25%; <u>and</u> identification of all first and last eluters of the eight homologue retentention time windows and documentation by labeling (F/L) on the chromatogram; <u>and</u> absolute retention times for switching from one homologous series to the next ≥ 10 seconds for all components of the mixture.	1) Readjust windows. 2) Evaluate system. 3) Perform maintenance. 4) Reanalyze CPSM. 5) No corrective action is necessary if 2,3,7,8-TCDD is not detected and the % valley is greater than 25%.		
GC/HRMS	ICAL = Minimum five-point initial calibration for target analytes, lowest concentration standard at or near the reporting limit.	ICAL prior to sample analysis, as needed by the failure of calibration verification, and when a new lot is used as a standard source for calibration verification, internal standard or recovery standard solutions.	RSD ≤ 20% for response factors for 17 unlabeled isomers & 9 labeled IS, <u>and</u> ion abundance ratios within limits specified in SOP; <u>and</u> S/N ≥ 10:1for target analytes.	Evaluate standard and instrument response. If problem with instrument (autosampler failure, response poor, etc.) or standards, correct as appropriate, then repeat initial calibration.		
GC/HRMS	ICV	Immediately following ICAL.	All project analytes must be within ± 30% of the expected value from the ICAL.	Evaluate standards and instrument response. If standard issue, repeat or remake then repeat standard as appropriate. If still fails, repeat initial calibration	Lab Manager / Analyst	WS-ID-0005

Instrument	Calibration Procedure	Frequency of Calibration	Acceptance Criteria	Corrective Action	Person Responsible for Corrective Action	SOP Reference
GC/HRMS	CCV	At the beginning of each 12-hour period, and at the end of each analytical sequence.	Ion abundance ratios must be in accordance with SOP; <u>and</u> RF (unlabeled standards) within ± 20%D of average RF from ICAL; <u>and</u> RF (labeled standards) within ± 30%D of average RF from ICAL.	Correct problem, repeat calibration verification. If fails, repeat ICAL and reanalyze all samples analyzed since last successful CCV End of Run CCV: If RF (unlabeled standards) > ± 20%D and ≤ ± 25%D and/or RF (labeled standards) > ± 30%D and ≤ ± 35%D of the average RF from ICAL use mean RF from bracketing CCVs to quantitate impacted samples. If bracketing CCVs differ by more than 25% RPD (unlabeled) or 35% RPD (unlabeled), run a new ICAL within 2 hours, and requantitate samples. Otherwise, reanalyze samples with positive detections.	Lab Manager / Analyst	WS-ID-0005
GC/FID ICAL ExTPH	ICAL	Instrument receipt, major instrument change, when CV does not meet criteria	One of the options below must be met:  Option 1: RSD for each analyte ≤ 20%;  Option 2: linear least squares regression: r ≥ 0.995;  Option 3: non-linear regression: COD r2 ≥ 0.99 (6 points shall be used for second order).	<ul><li>(1) Perform instrument maintenance as needed.</li><li>(2) Reanalyze and or reprep calibration standards.</li></ul>	Analyst, Department Manager	CA-315
	ICV	Immediately following ICAL.	All project analytes must be within established retention time windows. All project analytes must be within ± 20% of expected value from the ICAL.	Correct problem, rerun ICV. If that fails, repeat ICAL.	Analyst, Department Manager	

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

Instrument	Calibration Procedure	Frequency of Calibration	Acceptance Criteria	Corrective Action	Person Responsible for Corrective Action	SOP Reference
	CCV	Daily prior to sample analysis and at intervals of not less than once every 20 samples. Also, at the end of the analysis sequence.	%D must be ≤ +/- 20%	(1) Evaluate the samples: If the %D>+20% and sample results are <pql, %d="" if="" narrate.="">±20% and is likely a result of matrix interference, narrate. All samples must be reanalyzed that fall within the standard that exceeded criteria and the last standard that was acceptable.</pql,>	Analyst, Department Manager	
GC/FID GRO	ICAL	Minimum 5 point calibration using a gasoline component mixture.	One of the options below must be met: Option 1: RSD for each analyte ≤ 20%; Option 2: linear least squares regression: r ≥ 0.995; Option 3: non-linear regression: COD r2 ≥ 0.99 (6 points shall be used for second order).	Repeat initial calibration.	Analyst, Department Manager	CA-316
	ICV	Immediately following ICAL.	All project analytes within established retention time windows. All project analytes within ± 20% of expected value from the ICAL.	Correct problem, rerun ICV. If that fails, repeat ICAL.	Analyst, Department Manager	
	CCV	Daily prior to sample analysis and at intervals of not less than once every 20 samples. Also, at the end of the analysis sequence.	%D must be ≤ +/- 20%.	Evaluate the samples – if the %D>+20% and sample results are <pql, %d="" if="" narrate.="">   result of matrix interference, narrate. All samples must be reanalyzed that fall within the standard that exceeded criteria and the last standard that was acceptable</pql,>		

- Refer to the Analytical SOP References table (Worksheet #23).
   For data validation, the criterion for ICSA will be < LOQ.</li>

# SAP Worksheet #25 -- Analytical Instrument and Equipment Maintenance, Testing, and Inspection Table (UFP-QAPP Manual Section 3.2.3)

Instrument / Equipment	Maintenance Activity	Testing Activity	Inspection Activity	Frequency	Acceptance Criteria	CA	Responsible Person <sup>2</sup>	SOP Reference
GC/ECD	Check pressure and gas supply daily. Change septa and/or liner as needed, replace or cut column as needed. Other maintenance specified in lab Equipment Maintenance SOP.	EDB/DBCP and PCBs	Injector liner, septa, column, column flow.	Prior to ICAL and/or as necessary.	Acceptable ICAL or CCV	Correct the problem and repeat ICAL or CCV.	Analyst, Department Manager	CA-319, CA-329
GC/MS	Check pressure and gas supply daily. Manual tune if DFTPP not in criteria, change septa as needed, change liner as needed, cut column as needed. Other maintenance specified in lab Equipment Maintenance SOP	PAHs	Ion source, injector liner, column, column flow	Prior to ICAL and/or as necessary.	Acceptable ICAL or CCV	Correct the problem and repeat ICAL or CCV.	Analyst, Department Manager	CA-213
ICP-MS	Clean torch assembly and spray chamber when discolored or when degradation in data quality is observed. Clean nebulizer, check argon, replace peristaltic pump tubing as needed. Other maintenance specified in lab Equipment Maintenance SOP.	Metals	Torch, nebulizer, spray chamber, pump tubing	Prior to ICAL and as necessary	Acceptable ICAL or CCV	Correct the problem and repeat ICAL or CCV.	Analyst, Department Manager	CA-627
Mercury Analyzer	Replace peristaltic pump tubing, replace mercury lamp, replace drying tube, clean optical cell and/or clean liquid/gas separator as needed. Other maintenance specified in lab Equipment Maintenance SOP.	Mercury	Tubing, sample probe, optical cell	Prior to ICAL and as necessary	Acceptable ICAL or CCV	Correct the problem and repeat ICAL or CCV.	Analyst, Department Manager	CA-611, CA-615
GC/HRMS	Parameter Setup	Dioxins/ Physical check	Physical check	Initially; prior to DCC	Correct Parameters	Reset if incorrect	TestAmerica Chemist	WS-ID- 0005
GC/HRMS	Tune Check	Dioxins/ Instrument Performance	Conformance to instrument tuning.	Initially; prior to DCC	Compliance to ion abundance criteria	Correct the problem and repeat tune check	TestAmerica Chemist	WS-ID- 0005

Instrument / Equipment	Maintenance Activity	Testing Activity	Inspection Activity	Frequency	Acceptance Criteria	CA	Responsible Person <sup>2</sup>	SOP Reference
GC/FID	Check pressure and gas supply daily. Change septa and/or GC injector glass liner as needed. Replace or cut GC column as needed. Other maintenance specified in lab Equipment Maintenance SOP.	ExTPH	Injector liner, septa, column, column flow.	Prior to ICAL and/or as necessary.	Acceptable ICAL or CCV	Correct the problem and repeat ICAL or CCV.	Analyst, Department Manager	CA-315
GC/FID	Change septa, and/or GC injector glass liner as needed. Replace or cut GC column as needed. Bake out trap and column, change trap as needed.	GRO	Injector liner, septa, column, column flow.	Prior to ICAL and/or as necessary.	Acceptable ICAL or CCV	Correct the problem and repeat ICAL or CCV.	Analyst, Department Manager	CA-316

### **Project-Specific Sampling and Analysis Plan**

Site Name: Tank Farm 2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

## SAP Worksheet #26a – Sample Handling System

(UFP-QAPP Manual Appendix A)

## Sample Handling System - Katahdin

SAMPLE COLLECTION.	PACKAGING	AND SHIPMENT

Sample Collection (Personnel/Organization): FOL, Tetra Tech

Sample Packaging (Personnel/Organization): FOL, Tetra Tech

Coordination of Shipment (Personnel/Organization): FOL, Tetra Tech.

Type of Shipment/Carrier: Hand carrier or overnight courier service (Federal Express)

### SAMPLE RECEIPT AND ANALYSIS

Sample Receipt: Sample Custodians / Katahdin

Sample Custody and Storage: Sample Custodians / Katahdin

Sample Preparation: Extraction Lab, Metals Preparation Lab / Katahdin

Sample Determinative Analysis: Gas Chromatography Lab, Gas Chromatography/Mass Spectrometry Lab, Metals Lab / Katahdin

### SAMPLE ARCHIVING

Field Sample Storage (No. of days from sample collection): 60 days from receipt

Sample Extract/Digestate Storage (No. of days from extraction/digestion): 3 months from sample digestion/extraction

Biological Sample Storage (No. of days from sample collection): N/A

### SAMPLE DISPOSAL

Personnel/Organization: Sample Custodians/ Katahdin

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

# SAP Worksheet #26b – Sample Handling System – [Lab] (UFP-QAPP Manual Appendix A)

## Sample Handling System - Test America

Sample Handling System – Test America
SAMPLE COLLECTION, PACKAGING, AND SHIPMENT
Sample Collection (Personnel/Organization): Field Operation Leader, Tetra Tech
Sample Packaging (Personnel/Organization): Field Operation Leader, Tetra Tech
Coordination of Shipment (Personnel/Organization): Field Operation Leader, Tetra Tech
Type of Shipment/Carrier: Overnight courier service (FedEx)
SAMPLE RECEIPT AND ANALYSIS
Sample Receipt (Personnel/Organization): Sample Custodians, TestAmerica
Sample Custody and Storage (Personnel/Organization): Sample Custodians, TestAmerica
Sample Preparation (Personnel/Organization): Sample Preparation Technicians, TestAmerica
Sample Determinative Analysis (Personnel/Organization): Sample Analysis Technicians, TestAmerica
SAMPLE ARCHIVING
Field Sample Storage (No. of days from sample collection): 60 days
Sample Extract/Digestate Storage (No. of days from extraction/digestion): 60 days
Biological Sample Storage (No. of days from sample collection): Not applicable
SAMPLE DISPOSAL (Intact leftover samples)
Personnel/Organization: Sample Custodians, TestAmerica

Site Location: Portsmouth, Rhode Island

SAP Worksheet #27 - Sample Custody Requirements Table

(UFP-QAPP Manual Section 3.3.3)

Sample Designation and Tracking System

Each sample collected will be assigned a unique sample tracking number used to catalog the results. The

sample location IDs are listed in Worksheet #18. The sample tracking number for soil samples will

consist of alpha-numeric characters identifying the site, area, sample medium, location, and depth or

date. Any other pertinent information regarding sample identification will be recorded on the sample log

sheets, chain-of-custody forms, or in the field logbooks.

The alpha-numeric (A-N) coding to be used in the sample system is detailed below and in the subsequent

definitions.

AAA - NN - AA-NNNN - NNNN

(Site ID) - (Tank or AOC ID) - (Medium & Location) - (Depth or Date)

Site identifier: "TF2" for Tank Farm 2

"AOC" for locations that were formerly investigated under TtEC as Areas of Concern.

B219 for locations at Building 219

BSA for locations at former buoy storage area

AOC ID: TF2-###, as was identified by TtEC (Appendix A-1).

Medium identifier:

"W" for aqueous blanks

"SB" for soil boring samples

"SS" for surface soil samples

Sample location identifier: Each sample station is assigned a unique location identifier composed of

sequential numeric characters as shown on Worksheet #18.

Depth/Date:

For soil sample locations, this portion of the sample tracking number will represent the depth in feet bgs

from which the sample was collected; e.g., for soil samples collected from 2 to 4 feet bgs, this portion of

the sample tracking number will be "0204".

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**Project-Specific Sampling and Analysis Plan** 

Site Name: Tank Farm 2, NAVSTA Newport Project Name: Data Gaps Assessment

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For example, a soil sample at the Site, from the 8 to 10 foot interval of soil boring SB1060 will be labeled

TF2-SB1060-0810.

Quality Control Samples (QC) samples collected during the investigation will use the same coding system

as for the environmental samples. Field QC sample types are presented in Worksheet #20. Field QC

designations will conform to the following formats:

Field Duplicates: Blind field duplicate samples will be designated such that the location designation

will be replaced with "DUP" followed by a sequential value (the nth duplicate sample collected during

that sampling event) and a chronological value (MMYY). The sample log sheet will note from which

sample location the duplicate was collected. For example, the first soil field duplicate sample

collected in August 2013 at the Site will be labeled TF2-SB-DUP01-0813 and the second field

duplicate collected in August will be TF2-SB-DUP02-0813.

Rinsate Blanks: Rinsate blank sample identifiers will consist of the site, the medium (with "W" instead

of "MW", the "RB" label, and the date (MMYY). Example: TF2-W-RB01-0813.

Trip Blanks: will consist of the site, the medium (with "W" instead of "MW"), the "TB" label and the

date (MMYY).

Laboratory QC samples (matrix spike and laboratory duplicate samples) have no separate sample

identifier codes, but are noted on the chain-of-custody record and sample log sheet.

Sample Handling and Chain-of Custody Procedures

Custody of samples must be maintained and documented at all times. To ensure the integrity of a

sample from collection through analysis, an accurate written record is necessary to trace the possession

and handling of the sample. This documentation is referred to as the "chain of custody" form. Chain of

custody begins when samples are collected in the field, and is maintained by storing the samples in

secure areas until custody can be passed on. All samples will be accompanied by a chain-of-custody

form that will describe the sample identifiers, the analytical parameters, and the persons who are

responsible for the sample integrity.

Following collection, samples will be placed on ice in a secure cooler and attended by Tetra Tech

personnel or placed in locked vehicles or designated storage areas until analysis or shipment to an off-

site laboratory. Chain of custody procedures are described in further detail in the following Tetra Tech

SOPs:

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**Project-Specific Sampling and Analysis Plan** 

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SA-6.3 Field Documentation

SA-6.1 Non-Radiological Sample Handling

The samples will be shipped to the laboratories in coolers packed with ice and bubble wrap, or equivalent packing material, to cushion the samples to prevent breakage and to maintain the required temperature for the samples. A container filled with water and labeled "temperature blank" will be included in each cooler. The temperature of this blank will be measured by the laboratory upon sample receipt to verify acceptable sample preservation temperature. The coolers will be taped and sealed with a signed custody seal to ensure the chain of custody is maintained. The chain-of-custody forms are shipped to the laboratory with the samples.

Samples will be shipped to the laboratories by an overnight courier to ensure that maximum sample holding times are not exceeded. The maximum allowable sample holding times before sample extraction, digestion, or analysis are presented in Worksheet #19. Saturday deliveries will be coordinated by the FOL or his or her designee with the laboratory. Worksheet #19 also lists the sample containers, chemical preservatives, and temperature condition requirements to maintain the integrity of the sample.

The field crew will attempt to identify any potentially high concentration samples on the chain-of-custody form.

Laboratory procedures for sample receiving and chain-of-custody are detailed in SOP SD-902 (Katahdin) and SOP WS-QA-0003, (TestAmerica); and the laboratory procedures for disposal of the environmental samples are described in SOP 903 (Katahdin) and SOP WS-EHS-0001, (TestAmerica). These SOPs are included in Appendix G.

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

# Worksheet #28a – Laboratory QC Samples Table (UFP-QAPP Manual Section 3.4)

Note: Katahdin's statistically-derived QC limits referenced in the worksheets below refer to Katahdin's limits at the time of analysis. Katahdin's current limits are presented in Appendix G.

Matrix	Water / Soil					
Analytical Group	PAHs					
Analytical Method/ SOP Reference	SW-846 8270D SIM / CA-213					
QC Sample:	Frequency/ Number	Method/SOP QC Acceptance Limits	Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria (MPC)
Method Blank	One per preparation batch of twenty or fewer samples of similar matrix.	No target compounds > ½ LOQ (> LOQ for common laboratory contaminants) and > 1/10 the amount measured in any sample or 1/10 the PSL, whichever is greater.	Correct the problem. Report sample results that are <lod or="">10x the blank concentration. Reprepare and reanalyze the method blank and all associated samples with results &gt; LOD and &lt; 10x the contaminated blank result.</lod>	Analyst, Laboratory Department Manager and Data Validator	Bias/contamination	Same as Method/SOP QC Acceptance Limits.
Surrogate	3 per sample 2- Methylnaphthalene -d10 Fluorene-d10 Pyrene-d10	%R must be within Katahdin's statistically-derived QC limits.		Analyst, Laboratory Department Manager, and Data Validator	Accuracy/Bias	Same as Method/SOP QC Acceptance Limits.
LCS	One per preparation batch of twenty or fewer samples of similar matrix.	%R must be within Katahdin's statistically-derived QC limits, allowing for the number of	Correct problem, then reprepare and reanalyze the LCS and all samples in the associated preparatory batch for failed analytes, if sufficient sample material is available (see full explanation in Appendix F).	Analyst, Laboratory Department Manager, and Data Validator	Accuracy / Bias	Same as Method/SOP QC Acceptance Limits.

Matrix	Water / Soil					
Analytical Group	PAHs					
Analytical Method/ SOP Reference	SW-846 8270D SIM / CA-213					
QC Sample:	Frequency/ Number	Method/SOP QC Acceptance Limits	Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria (MPC)
		marginal exceedances presented in DoD QSM Table G-1.	Contact Client if samples cannot be reprepared within hold time.			
MS/MSD (not applicable for rinsate blanks)	One per sample delivery group (SDG) or every 20 samples.	%R should be within Katahdin's statistically-derived QC limits.  Water Precision: RPD should be ≤ 30%.  Soil Precision: RPD should be ≤ 50%.	Corrective actions will not be taken for samples when recoveries are outside limits if likely due to matrix, otherwise contact client.	Analyst, Laboratory Department Manager, and Data Validator	Precision/Accuracy/ Bias	Same as Method/SOP QC Acceptance Limits.
IS	Six per sample – 1,4- Dichlorobenzene- d4 Naphthalene-d8 Acenaphthene-d10 Phenanthrene-d10 Chrysene-d12 Perylene-d12	Retention times for internal standards must be ± 30 seconds and the responses within - 50% to +100% of the ICAL midpoint.	Inspect mass spectrometer or gas chromatograph for malfunctions. Mandatory reanalysis of samples analyzed while system was malfunctioning.	Analyst, Laboratory Department Manager, and Data Validator	Accuracy/ Bias	Same as Method/SOP QC Acceptance Limits.

Matrix	Soil/Water (rinsate	blanks)				
Analytical Group	PCBs					
Analytical Method/SOP Reference	SOP SW846 8082A /CA-329					
QC Sample:	Frequency/ Number	Method/SOP QC Acceptance Limits	Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria
Method Blank	One per preparation batch of 20 or fewer samples of similar matrix.	Contaminants in the method blank must be < ½ LOQ,	Correct the problem. Report sample results that are <lod or="">10x the blank concentration. Re-prepare and reanalyze the method blank and all associated samples with results &gt; LOD and &lt; 10x the contaminated blank result. Contact Client if samples cannot be re-prepared within hold time.</lod>	Analyst, Laboratory Department Manager and Data Validator Analyst, Laboratory Department Manager	Bias/contaminat ion	Same as Method/SOP QC Acceptance Limits.
Surrogates	PCBs: one per sample: Decachloro- biphenyl	%Rs must meet the laboratory statistically-derived control limits.	For QC and field samples, correct problem then re-prepare and reanalyze all failed samples for failed surrogates in the associated preparatory batch, if sufficient sample material is available. If obvious chromatographic interference with surrogate is present, reanalysis may not be necessary.  Contact Client if samples cannot be reprepared within hold time.	Analyst, Laboratory Department Manager	Accuracy/Bias	Same as Method/SOP QC Acceptance Limits.
LCS	One per preparation batch of 20 or fewer samples of similar matrix.	%R must be within Katahdin's statistically-derived QC limits.	Correct problem, then re-prepare and reanalyze the LCS and all samples in the associated preparatory batch for failed analytes, if sufficient sample material is available.  Contact Client if samples cannot be reprepared within hold time.	Analyst, Laboratory Department Manager	Accuracy/Bias	Same as Method/SOP QC Acceptance Limits.

Matrix	Soil/Water (rinsate	blanks)				
Analytical Group	PCBs					
Analytical Method/SOP Reference	SW846 8082A /CA	-329				
QC Sample:	Frequency/ Number	Method/SOP QC Acceptance Limits	Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria
MS/MSD (not applicable for rinsate blanks)	One per SDG or every 20 samples.	%R should be within Katahdin statistically derived limits. Soil Precision: RPD should be ≤ 50%. Water Precision: RPD should be ≤ 30%.	Corrective actions will not be taken for samples when recoveries are outside limits if likely due to matrix, otherwise contact client.	Analyst, Laboratory Department Manager	Precision/Accur acy/Bias	Same as Method/SOP QC Acceptance Limits.
Second Column Confirmation	All positive results must be confirmed.	Results between primary and second column must be RPD ≤ 40%. The higher of the two results will be reported unless matrix interference is apparent.	None. Apply qualifier if RPD >40% and discuss in the case narrative.	Analyst, Laboratory Department Manager	Precision	Same as Method/SOP QC Acceptance Limits.
Results between DL and LOQ	NA	Apply "J" qualifier to results between DL and LOQ.	NA	Analyst, Laboratory Department Manager	Accuracy	Same as QC Acceptance Limits.

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# Worksheet #28b – Laboratory QC Samples Table (UFP-QAPP Manual Section 3.4)

Matrix	Water / Soil					
Analytical Group	Metals (ICP-MS)					
Analytical Method/ SOP Reference	SW846 6020A / CA-627					
QC Sample:	Frequency/ Number	Method/SOP QC Acceptance Limits	Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria (MPC)
Method Blank	One per digestion batch of 20 or fewer samples of similar matrix.	No target metals > ½ LOQ and > 1/10 the amount measured in any sample or 1/10 the PSL, whichever is greater. For negative blanks, absolute value < LOD.	Correct the problem. Report sample results that are <lod or="">10x the blank concentration. Reprepare and reanalyze the method blank and all associated samples with results &gt; LOD and &lt; 10x the contaminated blank result.</lod>	Analyst, Laboratory Department Manager and Data Validator	Bias/contamination	Same as Method/SOP QC Acceptance Limits.
LCS	One per digestion batch of 20 or fewer samples of similar matrix.	Water and Soil: %R must be within 80-120%, allowing for the marginal exceedances presented in DoD QSM Table G-1.	Redigest and reanalyze all associated samples for affected analyte.	Analyst, Laboratory Department Manager, and Data Validator	Accuracy/Bias/ Contamination	Same as Method/SOP QC Acceptance Limits.
MS (not applicable for rinsate blanks)	One per sample delivery group (SDG) or every 20 samples.	%R should be within 80-120%if sample < 4x spike added.	Flag results for affected analytes for all associated samples with "N".	Analyst, Laboratory Department Manager, and Data Validator	Accuracy/Bias	%R should be within 75- 125%if sample < 4x spike added.
Post-digestion Spike (not applicable for rinsate blanks)	Project-specific frequency: When MS recovery fails or analyte concentration in all samples < 50x LOD	%R should be within 75-125%.	Run associated samples by method of standard addition or flag results.	Analyst, Laboratory Department Manager, and Data Validator	Accuracy/Bias	Same as Method/SOP QC Acceptance Limits.

Matrix	Water / Soil					
Analytical Group	Metals (ICP-MS)					
Analytical Method/ SOP Reference	SW846 6020A / CA-627					
QC Sample:	Frequency/ Number	Method/SOP QC Acceptance Limits	Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria (MPC)
Laboratory Duplicate (not applicable for rinsate blanks)	One per SDG or every 20 samples.	Project-specific criteria: If values are ≥ 5x LOQ, RPD should be ≤ 20%. If values are < 5x LOQ, Absolute Difference should be ≤ LOQ.	Flag results for affected analytes for all associated samples.	Analyst, Laboratory Department Manager, and Data Validator	Precision	Waters: If values are ≥ 5x LOQ, RPD should be ≤ 20%; if values are < 5x LOQ, Absolute Difference should be ≤ LOQ.  Soils: If values are ≥ 5x LOQ, RPD should be ≤ 35%; if values are < 5x LOQ, Absolute Difference should be ≤ 2x LOQ.
ICP Serial Dilution (not applicable for rinsate blanks)	One per preparation batch of twenty or fewer samples of similar matrix.	If original sample result is at least 50x LOQ, 5-fold dilution must agree within ± 10% of the original result.	Flag results for affected analytes for all associated samples with "E".	Analyst, Laboratory Department Manager, and Data Validator	Accuracy/Bias	Same as Method/SOP QC Acceptance Limits.
IS	Appropriate IS required for all analytes in all samples. Mass of IS must be <50 amu different from that of analyte	For each sample, IS intensity within 30-120% of that of initial calibration standard.	Reanalyze affected samples.	Analyst, Supervisor, QA Manager	Accuracy/Bias	Same as Method/SOP QC Acceptance Limits.

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# Worksheet #28c – Laboratory QC Samples Table (UFP-QAPP Manual Section 3.4)

Matrix	Water / Soil					
Analytical Group	Metals (Mercury)					
Analytical Method/ SOP Reference	SW-846 7470A, 7471B / CA-611, CA-615					
QC Sample:	Frequency/ Number	Method/SOP QC Acceptance Limits	Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria (MPC)
Method Blank	One per digestion batch of 20 or fewer samples of similar matrix.	No mercury > ½ LOQ and > 1/10 the amount measured in any sample or 1/10 the PSL, whichever is greater. For negative blanks, absolute value < LOD.	Correct the problem. Report sample results that are <lod or="">10x the blank concentration. Reprepare and reanalyze the method blank and all associated samples with results &gt; LOD and &lt; 10x the contaminated blank result.</lod>	Analyst, Laboratory Department Manager and Data Validator	Bias/contamination	Same as Method/SOP QC Acceptance Limits.
LCS	One per digestion batch of 20 or fewer samples of similar matrix.	Water and Soil: %R must be within 80-120%.	Redigest and reanalyze all associated samples for affected analyte.	Analyst, Laboratory Department Manager, and Data Validator	Accuracy/Bias/ Contamination	Same as Method/SOP QC Acceptance Limits.
MS (not applicable for rinsate blanks)	One per sample delivery group (SDG) or every 20 samples.	%R should be within 80-120% if sample < 4x spike added.	Flag results for affected analytes for all associated samples with "N".	Analyst, Laboratory Department Manager, and Data Validator	Accuracy/Bias	Same as Method/SOP QC Acceptance Limits.
Laboratory Duplicate (not applicable for rinsate blanks)	One per sample delivery group (SDG) or every 20 samples.	Project-specific criteria: If values are ≥ 5x LOQ, RPD should be ≤ 20%. If values are < 5x LOQ, Absolute Difference should be ≤ LOQ.	Flag results for affected analytes for all associated samples.	Analyst, Laboratory Department Manager, and Data Validator	Precision	Waters: If values are ≥ 5x LOQ, RPD should be ≤ 20%; if values are < 5x LOQ, Absolute Difference should be ≤ LOQ.  Soils: If values are ≥ 5x LOQ, RPD should be ≤ 35%; if values are < 5x LOQ, Absolute Difference should be ≤ 2x LOQ.

## **Project-Specific Sampling and Analysis Plan** Site Name: Tank Farm 2, NAVSTA Newport

Site Name: Tank Farm 2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

## SAP Worksheet #28d -- QC Samples Table

(UFP-QAPP Manual Section 3.4)

Note: TestAmerica's statistically-derived QC limits referenced in the worksheet below refer to TestAmerica's limits at the time of analysis. TestAmerica's current limits are presented in Appendix G.

Matrix	Water and Soil							
Analytical Group	Dioxins							
Analytical Method/ SOP Reference	SW-846 8290/ WS-ID-0005							
QC Sample:	Number Acceptance Lin				Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria
Method Blank	One per preparation batch	Project specific criteria, if available. Otherwise, no target analytes detected ≥ LOD or ≥ 20% of the associated regulatory limit or ≥ 5% of the sample result for the analyte, whichever is greater. (OCDD is considered a common laboratory contaminant and treated accordingly).	Verify instrument clean (evaluate calibration blank & samples prior to method blank), then reanalyze. Evaluate to determine if systematic issue within laboratory. Correct, then re-prepare and reanalyze the method blank and all samples processed with the contaminated blank in accordance with DoD QSM requirements.  "Totals" are not considered "target analytes" – no corrective action or flagging is necessary for "totals".	Analyst, Laboratory Department Manager, and Data Validator	contamination	Same as Method/SOP QC Acceptance Limits.		
Internal Standard Spike	Every field sample, standard and QC sample	%R for each IS in the original sample (prior to dilutions) must be within 40-135% for all 2378-substituted internal standards.	Correct problem, then reprepare and reanalyze the samples with failed IS.	Analyst, Laboratory Department Manager, and Data Validator	Accuracy / Bias	Same as Method/SOP QC Acceptance Limits.		
LCS	One per sample preparation batch	%R must be within TestAmerica's statistically-derived control limits.	Reanalyze LCS once. If acceptable, report. Otherwise, evaluate and reprepare and reanalyze the LCS and all samples in the associated preparation batch for failed analytes, if sufficient sample material is available.	Analyst, Laboratory Department Manager, and Data Validator	Accuracy/ Bias	Same as Method/SOP QC Acceptance Limits.		

Matrix	Water and Soil		1			
Analytical Group	Dioxins		1			
Analytical Method/ SOP Reference	SW-846 8290/ WS-ID-0005					
QC Sample:	Frequency/ Number	Method/SOP QC Acceptance Limits	Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria
MS/MSD	One MS/MSD per analytical/preparation batch	%R must be within TestAmerica's statistically-derived control limits; RPD must be ≤ 20%.	Identify problem; if not related to matrix interference, re-extract and reanalyze MS/MSD and all associated batch samples in accordance with DoD QSM requirements.	Analyst, Laboratory Department Manager, and Data Validator	Accuracy/ Bias/ Precision	Same as Method/SOP QC Acceptance Limits.

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

# **SAP Worksheet #28e – Laboratory QC Samples Table** (UFP-QAPP Manual Section 3.4)

Matrix	Water / Soil		]			
Analytical Group	GRO					
Analytical Method/ SOP Reference	SW-846 8015C / CA	<b>A-316</b>				
QC Sample:	Frequency/ Number	Method/SOP QC Acceptance Limits	Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria (MPC)
Method Blank	One per preparation batch of twenty or fewer samples of similar matrix.	No analytes detected >1/2 the LOQ and >1/10 the amount measured in any sample or 1/10 the regulatory limit (whichever is greater).	Investigate source of contamination. Evaluate the samples and associated QC: i.e., if the blank results are above the LOQ, report samples results which are < LOQ and >10X the blank. Otherwise, reprepare a blank and the remaining samples.	Analyst, Supervisor, QA Manager	Accuracy/Bias, Contamination	Same as Method/SOP QC Acceptance Limits.
Surrogates	One per sample – BFB	%R must be within Katahdin's statistically-derived QC limits.	For QC and field samples, correct problem then reprepare and reanalyze all failed samples for failed surrogates in the associated preparatory batch, if sufficient sample material is available. If obvious chromatographic interference with surrogate is present, reanalysis may not be necessary.  Contact Client if samples cannot be reprepared within hold time.	Analyst, Supervisor, QA Manager	Accuracy/Bias	Same as Method/SOP QC Acceptance Limits.
LCS	One per preparation batch of twenty or fewer samples of similar matrix.	%R must be within Katahdin's statistically-derived QC limits.	(1) Evaluate the samples and associated QC: i.e. If an MS/MSD was analyzed and acceptable, narrate. If an LCS/LCSD was performed and only one of the set was unacceptable, narrate. If the surrogate recoveries in the LCS are low but are acceptable in the blank and samples, narrate. If the LCS recovery is high but the sample results are < LOQ, narrate. Otherwise, reprep a blank and the remaining samples.	Analyst, Supervisor, QA Manager	Accuracy/Bias	Same as Method/SOP QC Acceptance Limits.

Matrix	Water / Soil					
Analytical Group	GRO					
Analytical Method/ SOP Reference	SW-846 8015C / CA	A-316				
QC Sample:	Frequency/ Number	Method/SOP QC Acceptance Limits	Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria (MPC)
MS/MSD (not applicable for rinsate blanks)	One per SDG or every 20 samples.	%R should be within Katahdin's statistically-derived QC limits.  Water and Soil Precision: RPD should be ≤ 30%.	Corrective actions will not be taken for samples when recoveries are outside limits if likely due to matrix, otherwise contact client.	Analyst, Supervisor, QA Manager	Precision/Accura cy/Bias	Same as Method/SOP QC Acceptance Limits.

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

## Worksheet #28f - Laboratory QC Samples Table

(UFP-QAPP Manual Section 3.4)

Matrix	Water / Soil					
Analytical Group	ExTPH					
Analytical Method/ SOP Reference	SW-846 8015C / 0	CA-315				
QC Sample:	Frequency/ Number	Method/SOP QC Acceptance Limits	Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria (MPC)
Method Blank	One per preparation batch of twenty or fewer samples of similar matrix.	No analytes detected >1/2 the LOQ and >1/10 the amount measured in any sample or 1/10 the regulatory limit (whichever is greater).	Investigate source of contamination. Evaluate the samples and associated QC: i.e., if the blank results are above the LOQ, report samples results which are < LOQ and >10X the blank. Otherwise, reprepare a blank and the remaining samples.	Analyst, Supervisor, QA Manager	Accuracy/Bias, Contamination	Same as Method/SOP QC Acceptance Limits.
Surrogates	One per sample  Ortho-terphenyl	%R must be within Katahdin's statistically-derived QC limits.	For QC and field samples, correct problem then reprepare and reanalyze all failed samples for failed surrogates in the associated preparatory batch, if sufficient sample material is available. If obvious chromatographic interference with surrogate is present, reanalysis may not be necessary.  Contact Client if samples cannot be reprepared within hold time.	Analyst, Supervisor, QA Manager	Accuracy/Bias	Same as Method/SOP QC Acceptance Limits.

Matrix	Water / Soil					
Analytical Group	ExTPH					
Analytical Method/ SOP Reference	SW-846 8015C / 0	CA-315				
QC Sample:	Frequency/ Number	Method/SOP QC Acceptance Limits	Corrective Action	Person(s) Responsible for Corrective Action	Data Quality Indicator (DQI)	Measurement Performance Criteria (MPC)
LCS	One per preparation batch of twenty or fewer samples of similar matrix.	Katahdin's statistically-	(1) Evaluate the samples and associated QC: i.e. If an MS/MSD was analyzed and acceptable, narrate. If an LCS/LCSD was performed and only one of the set was unacceptable, narrate. If the surrogate recoveries in the LCS are low but are acceptable in the blank and samples, narrate. If the LCS recovery is high but the sample results are < LOQ, narrate. Otherwise, reprepare a blank and the remaining samples.	Analyst, Supervisor, QA Manager	Accuracy/Bias	Same as Method/SOP QC Acceptance Limits.
MS/MSD (not applicable for rinsate blanks)	One per SDG or every 20 samples.	%R should be within Katahdin's statistically-derived QC limits.  Water and Soil Precision: RPD should be ≤ 30%.	Corrective actions will not be taken for samples when recoveries are outside limits if likely due to matrix, otherwise contact client.	Analyst, Supervisor, QA Manager	Precision/Accur acy/Bias	Same as Method/SOP QC Acceptance Limits.

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# SAP Worksheet #29 -- Project Documents and Records Table (UFP-QAPP Manual Section 3.5.1)

Document	Where Maintained
Field Documents Field Logbook Field Sample Forms Chain-of-Custody Records Air Bills Sampling Instrument Calibration Logs Sampling Notes Drilling Logs Photographs FMR Forms SAP HASP	Field documents will be maintained in the project file located in the Tetra Tech Wilmington, Massachusetts office.
Laboratory Documents and Records - in the form of analytical data package: Sample receipt/login form Sample storage records Sample preparation logs Equipment calibration logs Sample analysis run logs Reported results for standards, QC checks, and QC samples Data completeness checklists Telephone logs Extraction/clean-up records Raw data	Laboratory documents will be included in the hardcopy and electronic deliverables from the laboratory. Laboratory data deliverables will be maintained in the Tetra Tech Wilmington, Massachusetts project file and in long-term data package storage at a third-party professional document storage firm.  Electronic data results will be maintained in a database on a password protected Structured Query Language (SQL) server.
Assessment Findings Field Sampling Audit Checklist (if conducted) Analytical Audit Checklist (if conducted) Data Validation Memoranda (include tabulated data summary forms)	All assessment documents will be maintained in the Tetra Tech Wilmington, Massachusetts project file.
Reports Data Report	All versions of the Project Report and support documents (e.g., Data Validation Reports) will be stored in hard copy in the Tetra Tech Wilmington, Massachusetts project file and electronically in the server library.

Site Name: Tank Farm 2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

## SAP Worksheet #30 -- Analytical Services Table

(UFP-QAPP Manual Section 3.5.2.3)

Matrix	Analytical Group	Sample Locations/ID Number	Analytical Method	Data Package Turnaround Time	Laboratory / Organization (name and address, contact person and telephone number)	Backup Laboratory / Organization (name and address, contact person and telephone number)
	GRO and ExTPH		SW-846 8015C			
Soil and Aqueous (Aqueous QC	PAHs		SW-846 8270D SIM		Katahdin Analytical Services, Inc.	
Blanks)					600 Technology Way	
	Metals	See Worksheet #18	SW-846 6020A, 7470A, 7471B	21 days	Scarborough, Maine 04074 Contact: Ms. Jennifer Obrin	Not applicable
					(207) 874-2400	
Soil (aqueous rinsate blank)	PCBs		SW-846-8082A			
Soil and Aqueous QC Blanks	Dioxins	See Worksheet #18	SW-846 8290	21 days	TestAmerica 880 Riverside Parkway West Sacramento, CA 95605  Contact: Mr. Jill Kellmann 916-374-4402	Not applicable

Data package deliverables are detailed in the Analytical Technical Specifications included in Appendix F. Data packages will be provided as both hardcopy and portable document format (.PDF). Laboratories will provide a Naval Installation Restoration Information Solutions (NIRIS) compatible electronic data deliverable (EDD). Data packages will be Contract Laboratory Program (CLP)-equivalent (i.e., they will contain CLP-equivalent summary forms and raw data). Data will be stored by the analytical laboratories for seven years.

## **SAP Worksheet #31 -- Planned Project Assessments Table**

## (UFP-QAPP Manual Section 4.1.1)

Assessment Type	Frequency	Internal or External	Organization Performing Assessment	Person(s) Responsible for Performing Assessment (title and organizational affiliation)	Person(s) Responsible for Responding to Assessment Findings (title and organizational affiliation)	Person(s) Responsible for Identifying and Implementing Corrective Actions (CA) (title and organizational affiliation)	Person(s) Responsible for Monitoring Effectiveness of CA (title and organizational affiliation)
Laboratory System Audit <sup>1</sup>	Every 2 years	External	DoD ELAP Accrediting Body	DoD ELAP Accrediting Body Auditor	Katahdin QAM TestAmerica QAM	Katahdin QAM TestAmerica QAM	DoD ELAP Accrediting Body Auditor

<sup>1. [</sup>Laboratories] successfully completed the DoD's Environmental Laboratory Accreditation Program (ELAP) audit for all analytical methods. A copy of [Laboratory] DoD ELAP accreditations are included in Appendix G.

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

## SAP Worksheet #32 -- Assessment Findings and Corrective Action Responses

(UFP-QAPP Manual Section 4.1.2)

Assessment Type	Nature of Deficiencies Documentation	Individual(s) Notified of Findings (name, title, organization)	Timeframe of Notification	Nature of Corrective Action Response Documentation	Individual(s) Receiving Corrective Action Response (name, title, organization)	Timeframe for Response
Laboratory System Audit	Written audit report	QAM, Katahdin QAM, TestAmerica	Not specified by DoD ELAP	Letter	DoD ELAP Accrediting Body	Specified by DoD ELAP

## **Project-Specific Sampling and Analysis Plan** Site Name: Tank Farm 2, NAVSTA Newport

Site Name: Tank Farm 2, NAVSTA Newpor Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

## **SAP Worksheet #33 -- QA Management Reports Table**

(UFP QAPP Manual Section 4.2)

Type of Report	Frequency (daily, weekly monthly, quarterly, annually, etc.)	Projected Delivery Date(s)	Person(s) Responsible for Report Preparation (title and organizational affiliation)	Report Recipient(s) (title and organizational affiliation)
Data validation report	Per SDG	Within 3 weeks of receipt of laboratory data	Project Chemist, Tetra Tech	PM, Tetra Tech Tetra Tech project file
Major analysis problem identification (internal memorandum)	When persistent analysis problems are detected	Immediately upon detection of problem (same day)	QAM, Tetra Tech	PM (Tetra Tech), Program Manager (Tetra Tech), Tetra Tech project file
Project monthly progress report <sup>1</sup>	Monthly for duration of the project	Monthly	PM, Tetra Tech	Navy, project file
Laboratory QA Report	When significant plan deviations result from unanticipated circumstances	Immediately upon detection of problem (same day)	PM, Katahdin PM, TestAmerica	Tetra Tech project file

<sup>1.</sup> The monthly progress report is an update for the Navy RPM and contract office. The report includes information such as activities completed, an updated schedule, identification of outstanding issues, plans for the next period, and a financial narrative.

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

# SAP Worksheet #34 -- Verification (Step I) Process Table (UFP-QAPP Manual Section 5.2.1)

Verification Input	Description	Internal / External	Responsible for Verification (name, organization)
Chain-of-Custody Forms	The Tetra Tech FOL or designee will review and sign the chain-of-custody form to verify that all samples listed are included in the shipment to the laboratory and the sample information is accurate. The forms will be signed by the sampler and a copy will be retained for the project file, the Tetra Tech PM, and the Tetra Tech Data Validators.	Internal	Sampler and FOL, Tetra Tech
	The Laboratory Sample Custodian will review the sample shipment for completeness, integrity, and sign accepting the shipment. The Tetra Tech Data Validators will check that the chain-of-custody form was signed and dated by the Tetra Tech FOL or designee relinquishing the samples and also by the Laboratory Sample Custodian receiving the samples for analyses.	Internal/ External	Laboratory Sample Custodian, Katahdin and TestAmerica     Data Validators, Tetra Tech
SAP Sample Tables/ Chain-of-Custody Forms	Verify that all proposed samples listed in the SAP tables have been collected.	Internal	FOL or designee, Tetra Tech
Sample Log Sheets	Verify that information recorded in the log sheets is accurate and complete.	Internal	FOL or designee, Tetra Tech
Sample coordinates	Verify that actual sample locations are correct and in accordance with the SAP proposed locations. Document any discrepancies in the final report.	Internal	Tetra Tech, FOL or designee
SAP/ Field Logs/ Analytical Data Packages	Ensure that all sampling SOPs were followed. Verify that deviations have been documented and MPCs have been achieved. Particular attention should be given to verify that samples were correctly identified, that sampling location coordinates are accurate, and that documentation establishes an unbroken trail of documented chain-of-custody from sample collection to report generation. Verify that the correct sampling and analytical methods/SOPs were applied. Verify that the sampling plan was implemented and carried out as written and that any deviations are documented.	Internal	PM or designee, Tetra Tech
SAP/ Laboratory SOPs/ Raw Data/ Applicable Control Limits Tables	Ensure that all laboratory SOPs were followed. Verify that the correct analytical methods/SOPs were applied. Establish that all method QC samples were analyzed and in control as listed in the analytical SOPs. If method QA is significantly out of control, the Laboratory QAM will contact the Tetra Tech PM via telephone or e-mail for guidance prior to report preparation.	Internal	Laboratory QAM, Katahdin and TestAmerica

Verification Input	Description	Internal / External	Responsible for Verification (name, organization)
SAP/ Chain-of-Custody Forms	Check that field QC samples listed in Worksheet #20 were collected as required.	Internal	FOL or designee, Tetra Tech
Analytical Data Packages	All analytical data packages will be verified internally for completeness by the laboratory performing the work. The Laboratory QAM will sign the case narrative for each data package.	Internal	Laboratory QAM, Katahdin and TestAmerica
Electronic Data Deliverables (EDDs)/ Analytical Data Packages	Each EDD will be verified against the chain-of-custody and hard copy data package for accuracy and completeness. Laboratory analytical results will be verified and compared to the electronic analytical results for accuracy. Sample results will be evaluated for laboratory contamination and will be qualified for false positives using the laboratory method/preparation blank summaries. Positive results reported between the DL and the LOQ will be qualified as estimated. Extraneous laboratory qualifiers will be removed from the validation qualifier.	External	Data Validators, Tetra Tech
	Each data package will be verified for completeness by the Tetra Tech Data Validator. Missing information will be requested by the Tetra Tech Data Validator from the Laboratory PM.	External	Data Validators, Tetra Tech

SAP Worksheet #35 -- Validation (Steps IIa and IIb) Process Table (UFP-QAPP Manual Section 5.2.2) (Figure 37 UFP-QAPP Manual) (Table 9 UFP-QAPP Manual)

Step IIa / IIb	Validation Input	Description	Responsible for Validation (name, organization)	
Ila	Chain-of-Custody Forms	Custody - Ensure that the custody and integrity of the samples was maintained from collection to analysis and the custody records are complete and any deviations are recorded. Review that the samples were shipped and store at the required temperature and sample pH for chemically-preserved samples meet the requirements listed in Worksheet #19. Ensure that the analyses were performed within the holding times listed in Worksheet #19.	Project Chemist or Data Validators, Tetra Tech	
Ila/IIb	SAP/ Laboratory Data Packages/ EDDs	Ensure that the laboratory QC samples listed in Worksheet #28 were analyzed and that the MPCs listed in Worksheet #12 were met for all field samples and QC analyses. Check that specified field QC samples were collected and analyzed and that the analytical QC criteria set up for this project were met.	Project Chemist or Data Validators, Tetra Tech	
		Check the field sampling precision by calculating the RPD for field duplicate samples. Check the laboratory precision by reviewing the RPD or percent difference values from laboratory duplicate analyses; MS/MSDs; and LCS/LCSD, if available. Ensure compliance with the methods and project MPCs accuracy goals listed in Worksheet #12.		
		Check that the laboratory recorded the temperature at sample receipt and the pH of the chemically preserved samples to ensure sample integrity from sample collection to analysis.		
		Review the chain-of-custody forms generated in the field to ensure that the required analytical samples have been collected, appropriate sample identifications have been used, and correct analytical methods have been applied. The Tetra Tech Data Validator will verify that elements of the data package required for validation are present, and if not, the laboratory will be contacted and the missing information will be requested. Validation will be performed as per Worksheet #36. Check that all data have been transferred correctly and completely to the final Structured Query Language (SQL) database.		

## SAP Worksheet #35 – Validation (Steps IIa and IIb) Process Table (Continued)

Step IIa / IIb	Validation Input	Description	Responsible for Validation (name, organization)
IIb	SAP/ Laboratory Data Packages/ EDDs	QA/QC - Ensure that all QC samples specified in the SAP were collected and analyzed and that the associated results were within prescribed SAP acceptance limits. Ensure that QC samples and standards prescribed in analytical SOPs were analyzed and within the prescribed control limits. If any significant QC deviations occur, the Laboratory QAM shall have contacted the Tetra Tech PM.	Project Chemist or Data Validators, Tetra Tech
		Deviations - Summarize deviations from methods, procedures, or contracts in the Data Validation Report. Determine the impact of any deviation from sampling or analytical methods and SOPs requirements and matrix interferences effect on the analytical results. Qualify data results based on method or QC deviation and explain all the data qualifications. Print a copy of the project database qualified data depicting data qualifiers and data qualifiers codes that summarize the reason for data qualifications. Determine if the data met the MPCs and discuss the potential impact of any deviations on the technical usability of the data.	

## SAP Worksheet #36 -- Validation (Steps IIa and IIb) Summary Table (UFP-QAPP Manual Section 5.2.2.1)

Step IIa / IIb	Matrix	Analytical Group	Validation Criteria	Data Validator (title and organizational affiliation)
lla and llb	Soil and Aqueous	PAHs.	Tier II <sup>(1)</sup> data validation. Project-specific criteria for PAHs by SW-846 8270D SIM, are listed in Worksheets #12, #15, #19, #24, and #28. Region I EPA-NE Data Validation Functional Guidelines for Evaluating Environmental Analyses, Part II, December 1996 (USEPA, 1996) will be applied using these criteria.	Tetra Tech, Project Chemist (K. Carper) and staff chemists
lla and llb	Soil and Aqueous	PCBs	Tier II <sup>(1)</sup> data validation. Criteria for EDB by SW-846 8011 are listed in Worksheets #12, #15, #19, #24, and #28. Region I EPA-NE Data Validation Functional Guidelines for Evaluating Environmental Analyses, Part III, February 2004 (EPA, 2004) will be applied using these criteria.	Tetra Tech, Project Chemist (K. Carper and staff chemists
IIa and IIb	Soil and Aqueous	Dioxins	Tier II <sup>(1)</sup> data validation. Project-specific criteria for dioxins by SW-846 8290 are listed in Worksheets #12, #15, #19, #24, and #28. USEPA National Functional Guidelines for Chlorinated Dibenzo-p-Dioxins (CDDs) and Chlorinated Dibenzofurans (CDFs) Data Review, September 2005 (EPA, 2005b) will be applied using these criteria.	Tetra Tech, Project Chemist (K. Carper) and staff chemists
IIa and IIb	Soil and Aqueous	Metals	Tier II <sup>(1)</sup> data validation. Project-specific criteria for metals by SW-846 6020A/7470A/7471B are listed in Worksheets #12, #15, #19, #24, and #28. Region I EPA-NE Data Validation Functional Guidelines for Evaluating Environmental Analyses, Part IV, November 2008 (USEPA, 2008) will be applied using these criteria.	Tetra Tech, Project Chemist (K. Carper) and staff chemists

<sup>1 –</sup> As defined in the Region I EPA-NE Data Validation Functional Guidelines for Evaluating Environmental Analyses, Part I, Attachment B, "Region 1 Tiered Organic and Inorganic Data Validation Guidelines", July 1, 1993, Draft (USEPA, 1996).

**Project-Specific Sampling and Analysis Plan** 

Site Name: Tank Farm 2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

SAP Worksheet #37 -- Usability Assessment

(UFP-QAPP Manual Section 5.2.3)

## **Data Usability Assessment**

The usability of the data directly affects whether project objectives can be achieved. The following characteristics will be evaluated at a minimum. The results of these evaluations will be included in the project report. The characteristics will be evaluated for multiple concentration levels if the evaluator determines that this is necessary. To the extent required by the type of data being reviewed, the assessors will consult with other technically competent individuals to render sound technical assessments of these data characteristics:

<u>Completeness</u>: The FOL, acting on behalf of the Project Team, will determine whether deviations from the scheduled sample collection or analyses occurred. If they have occurred and the Tetra Tech PM determines that the deviations compromise the ability to meet project objectives she will consult with the Navy RPM and other project team members, as necessary (determined by the Navy RPM), to develop appropriate corrective actions.

<u>Precision</u>: The Project Chemist, or designee, acting on behalf of the Project Team, will determine whether precision goals for field duplicates and laboratory duplicates were met. This will be accomplished by comparing duplicate results to precision goals identified in <u>Worksheets #12</u> and #28. This will also include a comparison of field and laboratory precision with the expectation that laboratory duplicate results will be no less precise than field duplicate results. If the goals are not met, or data have been flagged as estimated (J qualifier), limitations on the use of the data will be described in the project report.

<u>Accuracy</u>: The Project Chemist, acting on behalf of the Project Team, will determine whether the accuracy/bias goals were met for project data. This will be accomplished by comparing percent recoveries of LCS, LCSD, MS, MSD, and surrogate compounds to accuracy goals identified in Worksheet #28. This assessment will include an evaluation of field and laboratory contamination; instrument calibration variability; and analyte recoveries for surrogates, matrix spike, matrix spike duplicate, and laboratory control samples. If the goals are not met, limitations on the use of the data will be described in the project report. Bias of the qualified results and a description of the impact of identified non-compliances on a specific data package or on the overall project data will be described in the project report.

Representativeness: A project scientist identified by the Tetra Tech PM, and acting on behalf of the Project Team, will determine whether the data are adequately representative of intended populations, both spatially and temporally. This will be accomplished by verifying that samples were collected and analyzed in accordance with this SAP, by reviewing spatial and temporal data variations, and by comparing these characteristics to expectations. The usability report will describe the representativeness of the data for each matrix and analytical fraction. This will not require quantitative comparisons unless professional judgment of the project scientist indicates that a quantitative analysis is required.

<u>Comparability</u>: The Project Manager or designee, acting on behalf of the Project Team, will determine whether the data generated under this project are sufficiently comparable to historical property data generated by different methods and for samples collected using different procedures and under different property conditions. This will be accomplished by comparing overall precision and bias among data sets for each matrix and analytical fraction. This will not require quantitative comparisons unless the Project Chemist indicates that such quantitative analysis is required.

<u>Sensitivity</u>: The Project Chemist, acting on behalf of the Project Team, will determine whether project sensitivity goals listed in <u>Worksheet #15</u> are achieved. The overall sensitivity and quantitation limits from multiple data sets for each matrix and analysis will be compared. If sensitivity goals are not achieved, the limitations on the data will be described.

## **Project-Specific Sampling and Analysis Plan**

Site Name: Tank Farm 2, NAVSTA Newport Project Name: Data Gaps Assessment Site Location: Portsmouth, Rhode Island

Title: Sampling and Analysis Plan Document Number: W5211722F Revision Number: 0 Revision Date: July 2013

## Describe the evaluative procedures used to assess overall measurement error associated with the project:

After completion of the data validation, the data and data quality will be reviewed to determine whether sufficient data of acceptable quality are available for decision making. In addition to the evaluations described above, a series of inspections and statistical analyses will be performed to estimate these characteristics. The statistical evaluations will include simple summary statistics for target analytes, such as maximum concentration, minimum concentration, number of samples exhibiting non-detected results, number of samples exhibiting positive results, and the proportion of samples with detected and non-detected results. The Project Team members identified by the Project Manager will assess whether the data collectively support the attainment of project objectives. They will consider whether any missing or rejected data have compromised the ability to make decisions or to make the decisions with the desired level of confidence. The data will be evaluated to determine whether missing or rejected data can be compensated by other data. Although rejected data will generally not be used, there may be reason to use them in a weight-of-evidence argument, especially when they supplement data that have not been rejected. If rejected data are used, their use will be supported by technically defensible rationales.

For statistical evaluations, Worksheet #11 describes how to treat non-detected values.

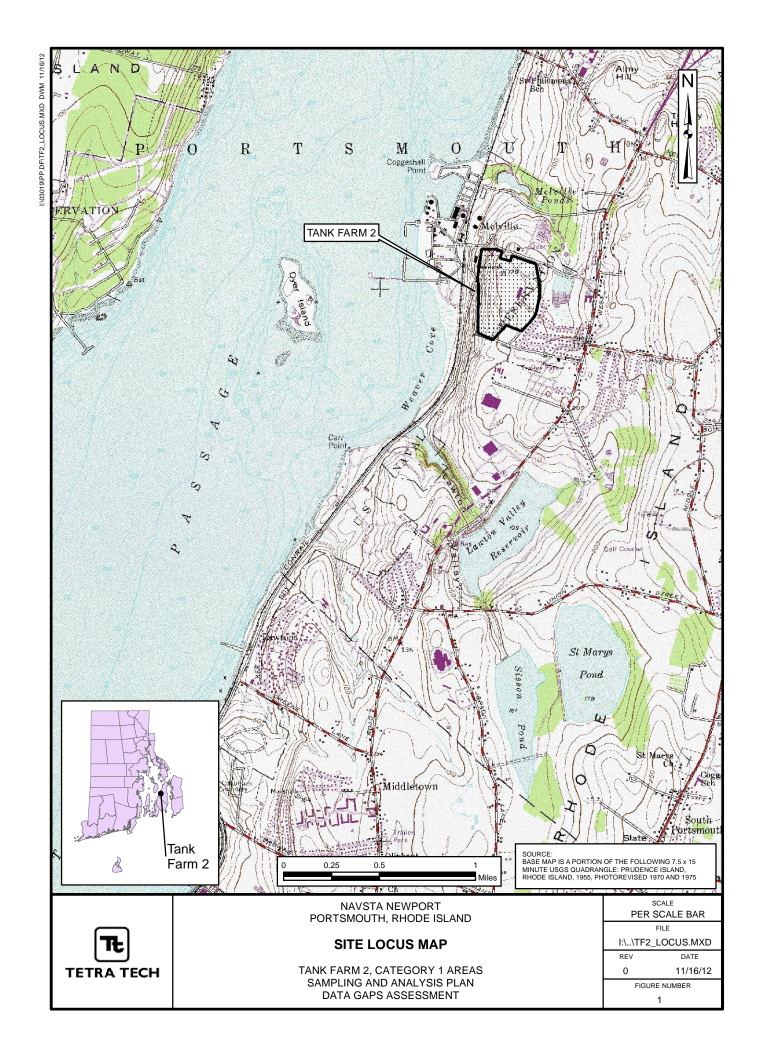
## Identify the personnel responsible for performing the usability assessment:

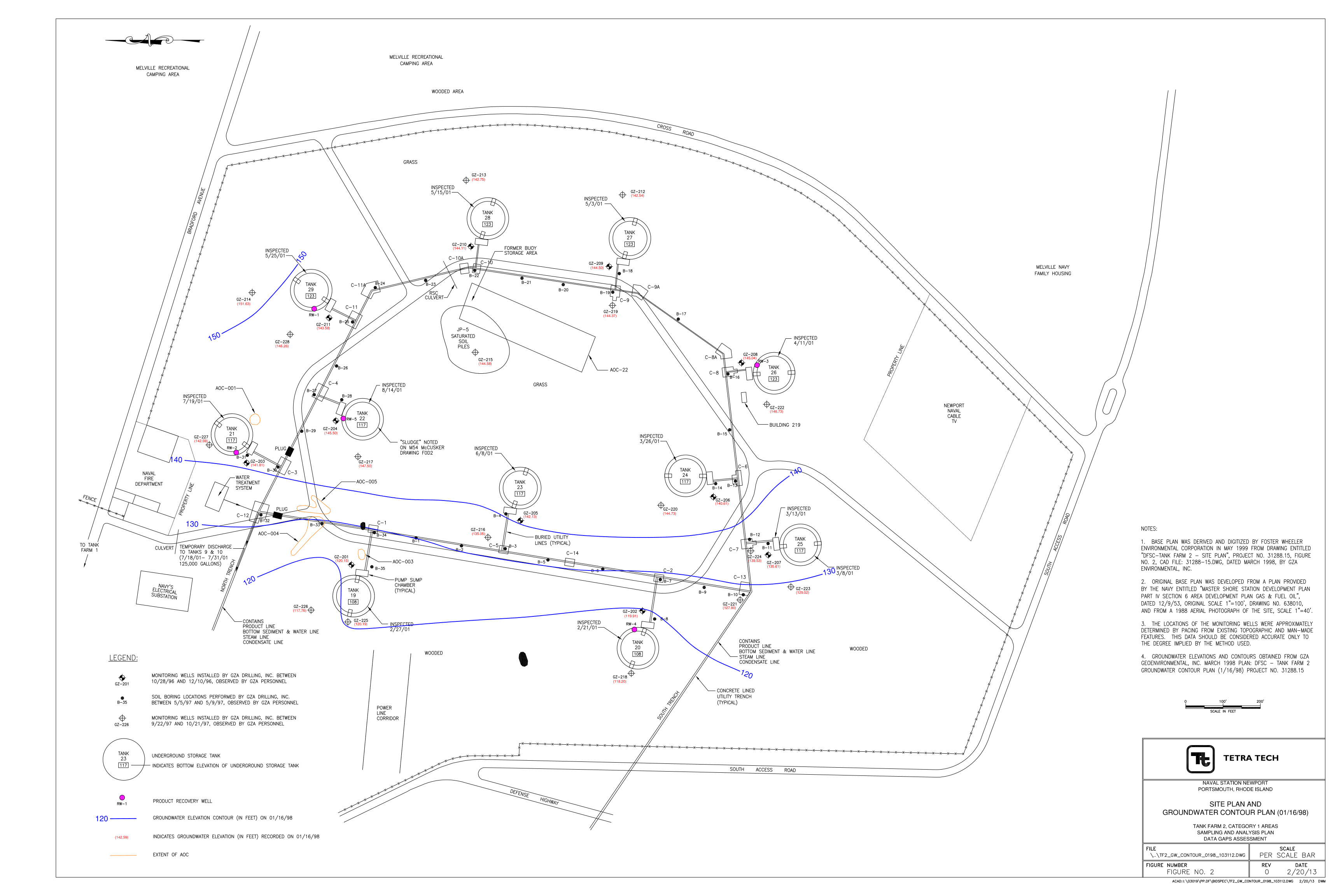
The Tetra Tech PM, Project Chemist, and FOL will be responsible for conducting the listed data usability assessments. The data usability assessment will be reviewed with the Project Team. If deficiencies affecting the attainment of project objectives are identified, the review will take place either in a face to face meeting or a teleconference depending on the extent of identified deficiencies. If no significant deficiencies are identified, the data usability assessment will simply be documented in the project report and reviewed during the normal document review cycle.

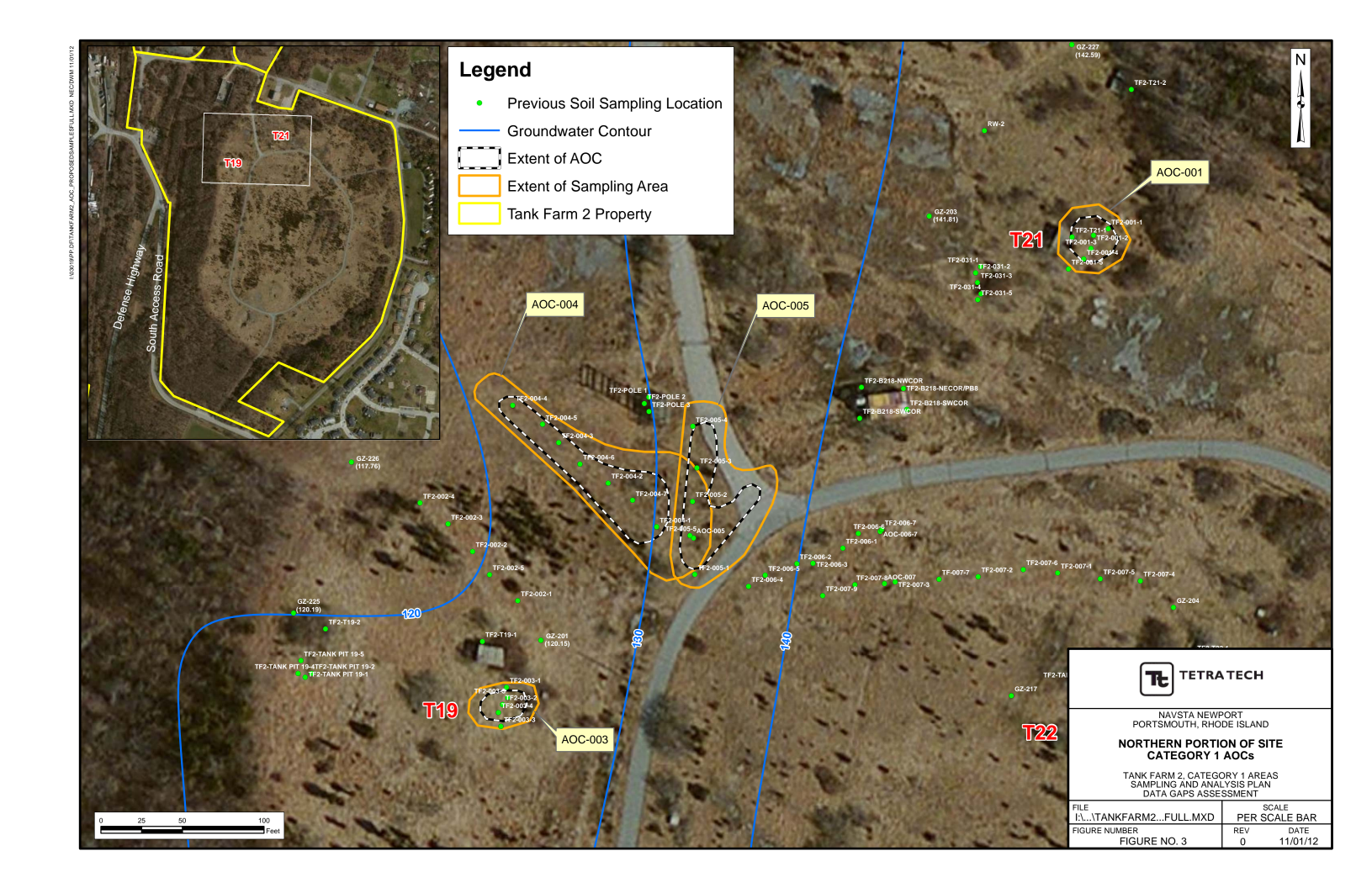
Describe the documentation that will be generated during usability assessment and how usability assessment results will be presented so that they identify trends, relationships (correlations), and anomalies:

The data will be presented in tabular format, including data qualifications such as estimation (J, UJ) or rejection (R). The project report will identify and describe the data usability limitations and suggest re-sampling or other corrective actions, if necessary. Graphical presentations of the data such as concentration tag maps will be generated as part of the overall data evaluation process.









RED TEXT/ARROWS = CONTAMINANT SOURCE/DISCHARGE ROUTE

BLUE TEXT/ARROWS = FATE AND TRANSPORT ROUTE

TETRA TECH

NAVAL STATION NEWPORT PORTSMOUTH, RHODE ISLAND

FIRST TIER CONCEPTUAL SITE MODEL

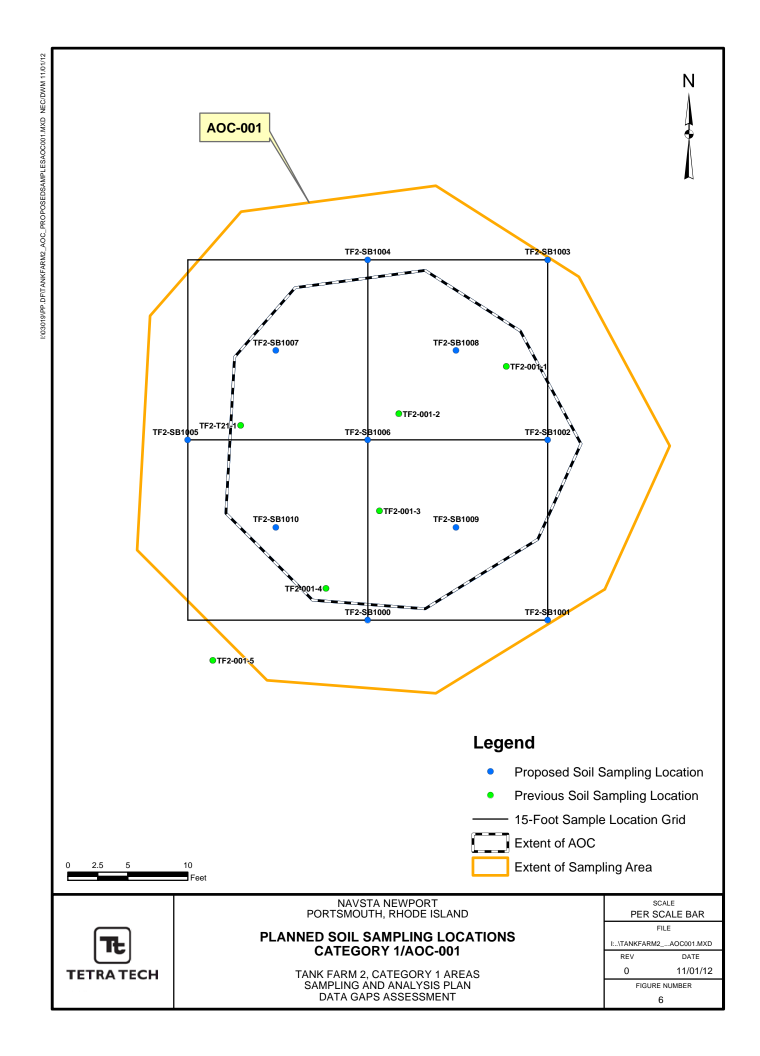
TANK FARM 2, CATEGORY 1 AREAS SAMPLING AND ANALYSIS PLAN DATA GAPS ASSESSMENT

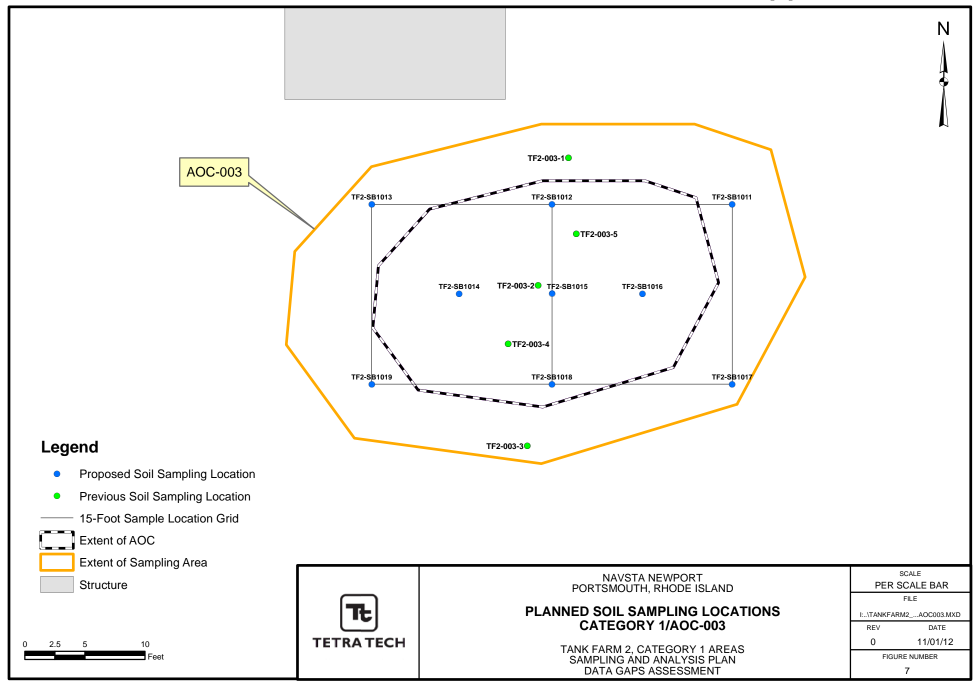
FILE \\TF2_CSM.DWG		scale TO SCALE
FIGURE NO. 4	<b>REV</b> O	<b>DATE</b> 11/1/12

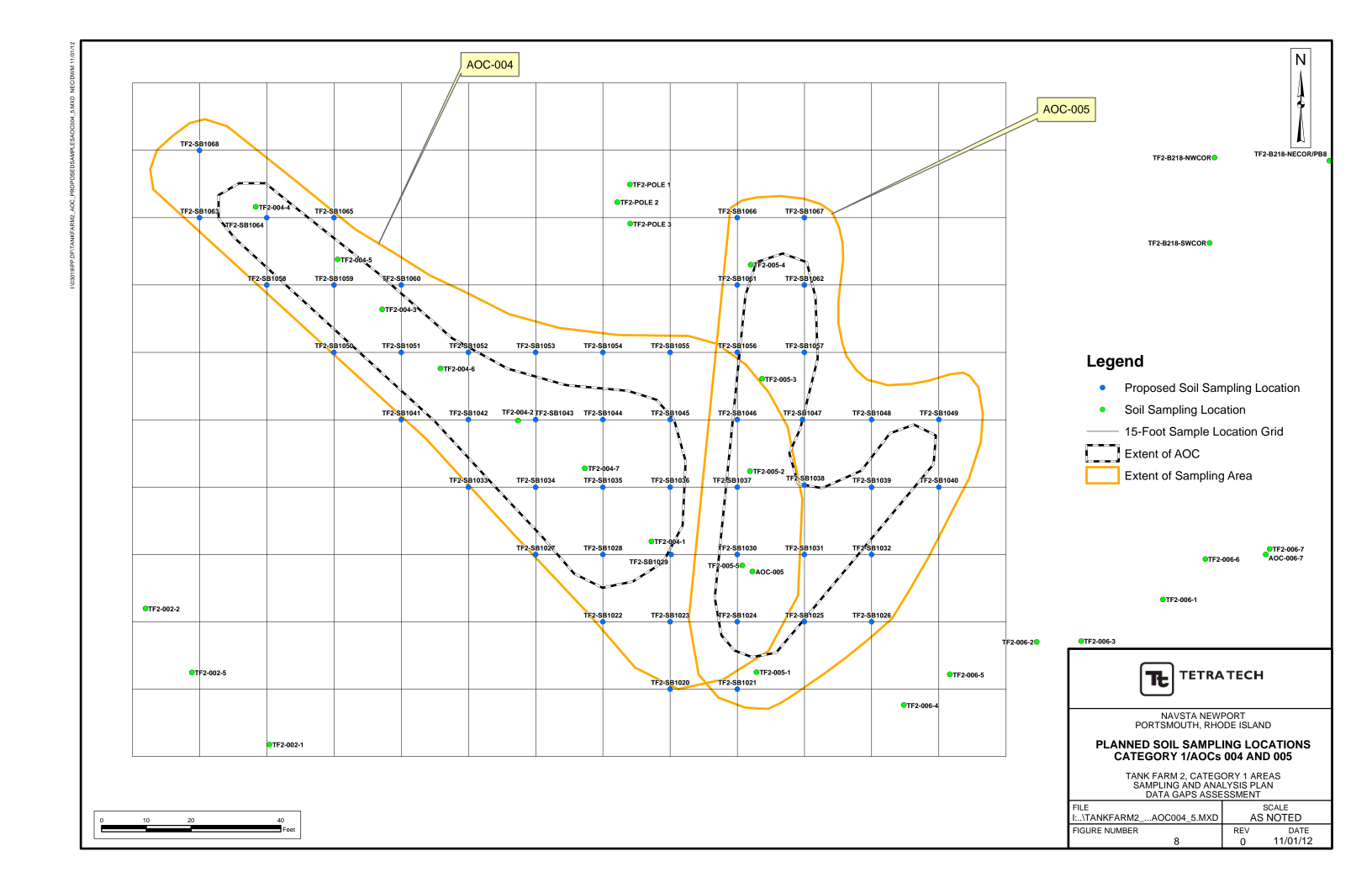
DATA GAPS ASSESSMENT

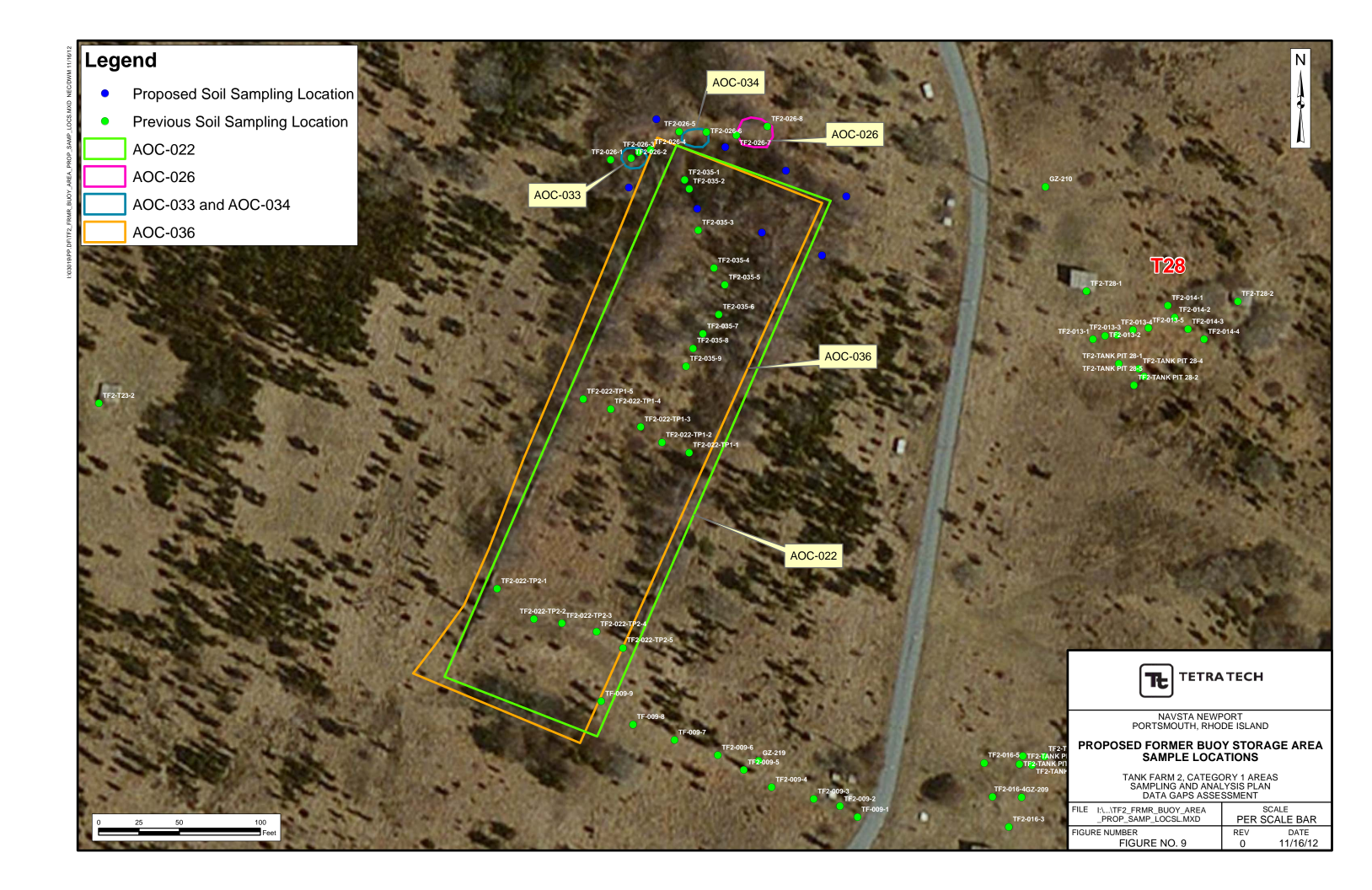
FIGURE NUMBER

FIGURE NO. 5











Site Name: Tank Farm 2 Project Name: NAVSTA Newport Site Location: Newport, Rhode Island Title: Data Gaps Assessment Document No.: W5211722F Revision Number: 0 Date: July 2013

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**Project-Specific Sampling and Analysis Plan** 

Site Name: Tank Farm 2 Project Name: NAVSTA Newport Site Location: Newport, Rhode Island Title: Data Gaps Assessment Document No.: W5211722F Revision Number: 0 Date: July 2013

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# APPENDIX A HISTORIC SITE CONDITIONS

# SUMMARY OF SITE CONDITIONS SAMPLING AND ANALYSIS PLAN

# REMEDIAL INVESTIGATION TANK FARM 2 NAVAL STATION, NEWPORT, NEWPORT, RHODE ISLAND

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Area	Description	History	Associated sampling locations	Available Data	Exceedances	Remedial Category	Other Information	Next Steps
Tank 19	2.5-million gallon capacity storage underground storage tank; stored No. 5 Fuel oil from the 1940's to 1975, distillate fuel from 1975 to 1985, and marine diesel fuel from 1985 until the mid 1990's.	infiltration points in the tanks. A marine chemist certified that the tanks were gas-free and safe for workers. Tanks	Borings: B-35   TtFC 2005 soil exploration: TF2-Tank19-1	Groundwater: TPH, VOC, SVOC, DRO Soil: PAH, TPH, VOC, Petroflag™ screening	None	2		No further action.
Tank 20	2.5-million gallon capacity storage underground storage tank; stored No. 5 Fuel oil from the 1940's to 1975, distillate fuel from 1975 to 1985, and marine diesel fuel from 1985 until the mid 1990's.	were cracks on the floor and the weeping of oil/water was observed. In 2001, FwEC pumped, cleaned, and repaired any	<b>TtEC 2005 soil exploration</b> : TF2-Tank20-1, TF2-Tank20-2, TF2-Tankpit20-1 through TF2-	Groundwater: TPH, VOC, SVOC, DRO, Oil-fingerprint Soil: PAH, TPH, VOC, Petroflag™ screening	<b>Groundwater:</b> Over 1-foot of LNAPL was detected in GZ-202 multiple times during gauging performed in 2009.	2		Conduct periodic gauging and bailing of LNAPL and install additional wells. Conduct a Site Investigation under RIDEM UST regulations, confirm petroleum is not mobile, develop a Corrective Action Plan accordingly.
Tank 21	2.5-million gallon capacity storage underground storage tank; stored No. 5 Fuel oil from the 1940's to 1975, distillate fuel from 1975 to 1985, and marine diesel fuel from 1985 until the mid 1990's.	were cracks on the floor and the weeping of oil/water was observed. In 2001, FwEC pumped, cleaned, and repaired any infiltration points in the tanks. A marine chemist certified	<b>TtEC 2005 soil exploration</b> : TF2-Tank21-1, TF2-Tank21-2, TF2-Tankpit21-1 through TF2-	Groundwater: TPH, VOC, SVOC, DRO, Lead Soil: DRO, PAH, TPH, VOC, Petroflag™ screening	None	2		No further action.
Tank 22	2.5-million gallon capacity storage underground storage tank; stored No. 5 Fuel oil from the 1940's to the 1970's. At that time, this tank was taken out of service, cleaned and used as a slop tank.	were cracks on the floor and the weeping of oil/water was observed. In 2001, FwEC pumped, cleaned, and repaired any	<b>TtEC 2005 soil exploration</b> : TF2-Tank22-1, TF2-Tank22-2, TF2-Tankpit22-1 through TF2-	<b>Groundwater</b> : PAH, TPH, VOC <b>Soil</b> : PAH, TPH, VOC, Petroflag™ screening	None	2		No further action.
Tank 23	2.5-million gallon capacity storage underground storage tank; stored No. 5 Fuel oil from the 1940's to 1975, distillate fuel from 1975 to 1985, and marine diesel fuel from 1985 until the mid 1990's.	infiltration points in the tanks. A marine chemist certified that the tanks were gas-free and safe for workers. Tanks	Monitoring Wells: GZ-205, GZ-216 Borings: B-4 TtEC 2005 soil exploration: TF2-Tank23-1, TF2-Tank23-2, TF2-Tankpit23-1 through TF2- Tankpit23-5	<b>Groundwater</b> : DRO, GRO, VOC, SVOC, Lead <b>Soil</b> : TPH, VOC, SVOC, Petroflag™ screening	None	2		No further action.

# SUMMARY OF SITE CONDITIONS SAMPLING AND ANALYSIS PLAN REMEDIAL INVESTIGATION TANK FARM 2

# NAVAL STATION, NEWPORT, NEWPORT, RHODE ISLAND PAGE 2 of 10

Area	Description	History	Associated sampling locations	Available Data	Exceedances	Remedial Category	Other Information	Next Steps
Tank 24	2.5-million gallon capacity storage underground storage tank; stored No. 5 Fuel oil from the 1940's to 1975, distillate fuel from 1975 to 1985, and marine diesel fuel from 1985 until the mid 1990's.	GZA cleaned and closed in 1996/1997. A structural assessment of the interior of the tank indicated that there were cracks on the floor and the weeping of oil/water was observed. In 2001, FwEC pumped, cleaned, and repaired any infiltration points in the tanks. A marine chemist certified that the tanks were gas-free and safe for workers. Tanks were ballasted with clean water. No closure permit has been received from RIDEM.	Monitoring Wells: GZ-206, GZ-220 Borings: B-13, B-14 TtEC 2005 soil exploration: TF2-Tank24-1, TF2-Tank24-2, TF2-Tankpit24-1 through TF2- Tankpit24-5	<b>Groundwater:</b> PAH, VOC, TPH <b>Soil</b> : PAH, TPH, VOC, Petroflag™ screening	None	2	GZ-206 could not be located in the TtEC 2009 GW sampling event.	No further action.
Tank 25	2.5-million gallon capacity storage underground storage tank; stored No. 5 Fuel oil from the 1940's to 1975, distillate fuel from 1975 to 1985, and marine diesel fuel from 1985 until the mid 1990's.	GZA cleaned and closed in 1996/1997. A structural assessment of the interior of the tank indicated that there were cracks on the floor. In 2001, FwEC pumped, cleaned, and repaired any infiltration points in the tanks. A marine chemist certified that the tanks were gas-free and safe for workers. Tanks were ballasted with clean water. No closure permit has been received from RIDEM.	Monitoring Wells: GZ-207, GZ-221, GZ-223, GZ-224 Borings: B-10, B-11, B-12 TtEC 2005 soil exploration: TF2-Tank25-1, TF2-Tank25-2, TF2-Tankpit25-1 through TF2-Tankpit25-6, TF2-Tankpit25-6D TtEC 2005 Remediation: TF2-T25-R1 through TF2-T25-R38	Groundwater: DRO, TPH, VOC, SVOC, Lead Soil: TPH, VOC, SVOC, DRO, Petroflag™ screening	No exceedences post-remedial excavation.	2	GZ-207 and GZ-224 were destroyed during the Tank 25 remediation in 2005 by TtEC.	No further action.
Tank 26	storage underground storage tank; stored No. 5 Fuel oil	GZA cleaned and closed in 1996/1997. A structural assessment of the interior of the tank indicated that there were cracks on the floor and the weeping of groundwater was observed. In 2001, FwEC pumped, cleaned, and repaired any infiltration points in the tanks. A marine chemist certified that the tanks were gas-free and safe for workers. Tanks were ballasted with clean water. No closure permit has been received from RIDEM.	TtEC 2005 soil exploration: TF2-Tank26-1, TF2-Tank26-2, TF2-Tankpit26-1 through TF2-	SVOC, Oil fingerprinting Soil: PAH, TPH, VOC,	<b>Groundwater:</b> LNAPL was detected in GZ-208 multiple times during 2009. The last time the well was gauged in 2009, LNAPL was greater than 1-foot thick.	2		Conduct periodic gauging and bailing of LNAPL and install additional wells. Conduct a Site Investigation under RIDEM UST regulations, confirm petroleum is not mobile, develop a Corrective Action Plan accordingly.
Tank 27	storage underground storage tank; stored No. 5 Fuel oil	GZA cleaned and closed in 1996/1997. A structural assessment of the interior of the tank indicated that there were cracks on the floor and the weeping of oil/water was observed. In 2001, FwEC pumped, cleaned, and repaired any infiltration points in the tanks. A marine chemist certified that the tanks were gas-free and safe for workers. Tanks were ballasted with clean water. No closure permit has been received from RIDEM.	Monitoring Wells: GZ-209, GW-212, GZ-219 Borings:B-18 TtEC 2005 soil exploration: TF2-Tank27-1, TF2-Tank27-2, TF2-Tankpit27-1 through TF2- Tankpit27-5	Groundwater: DRO, TPH, VOC, SVOC, Lead Soil: TPH, DRO, SVOC, VOC, Petroflag™ screening	<b>Soil:</b> A soil sample collected at 15-17' in boring GZ-209 exceeded the RIDEM GB Leachability Criteria with a concentration of TPH = 5,600 mg/kg in 1996	2	In 2009, TtEC recorded that the cover and pad of well GZ-209 was damaged and could not be removed to initiate sampling.	No further action (no exceedence in groundwater sample from GZ-209).
Tank 28	storage underground storage tank; stored No. 5 Fuel oil	GZA cleaned and closed in 1996/1997. A structural assessment of the interior of the tank indicated that there were cracks on the floor and the weeping of groundwater was observed. In 2001, FwEC pumped, cleaned, and repaired any infiltration points in the tanks. A marine chemist certified that the tanks were gas-free and safe for workers. Tanks were ballasted with clean water. No closure permit has been received from RIDEM.	TtEC 2005 soil exploration: IF2-Tank28-1, TF2-Tank28-2, TF2-Tankpit28-1 through TF2- Tanknit28-5	Groundwater: TPH, VOC, SVOC Soil: PAH, TPH, VOC, Petroflag™ screening	None	2	TtEC found an obstruction in well GZ-213 during the 2009 sampling event.	No further action.

#### SUMMARY OF SITE CONDITIONS SAMPLING AND ANALYSIS PLAN REMEDIAL INVESTIGATION TANK FARM 2

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Area	Description	History	Associated sampling locations	Available Data	Exceedances	Remedial Category	Other Information Next Steps
Tank 29	distillate fuel from 1975 to 1985, and marine diesel fuel from 1985 until the mid	GZA cleaned and closed in 1996/1997. A structural assessment of the interior of the tank indicated that there were cracks on the floor. In 2001, FwEC pumped, cleaned, and repaired any infiltration points in the tanks. A marine chemist certified that the tanks were gas-free and safe for workers. Tanks were ballasted with clean water. No closure permit has been received from RIDEM.	Monitoring Wells: GZ-211, GZ-214, GZ-228 Recovery Wells: RW-1 Borings: B-25 TtEC 2005 soil exploration: TF2-Tank29-1, TF2-Tank29-2, TF2-Tankpit29-1 through TF2- Tankpit29-5	SVOC Soil: DRO, PAH, TPH, VOC,	<b>Groundwater:</b> Product was observed coming through the sample tubing at GZ-211 during the 2009 sampling round, and thus the well was not sampled.	2	Conduct periodic gauging and bailing of LNAPL and install additional wells. Conduct a Site Investigation under RIDEM UST regulations, confirm petroleum is not mobile, develop a Corrective Action Plan accordingly.
Underground Distribution Lines	This piping connects the tanks to one another and to the fuel distribution area. The lines are located approximately 10-feet underground in concrete lined utility trenches.	During 1996/1997 tank closure activities, GZA also decommissioned the fuel distribution pipelines associated with each tank and the transfer pipe loop. The pipes were cleaned until a PID detected an internal atmosphere of <25 ppm and then grouted. GZA also performed asbestos abatement activities on encountered sections of the piping with asbestos containing insulation. In 2001, FwEC inspected the sections of piping decommissioned by GZA. If a length of piping was found to have elevated levels of VOCs, it was recleaned and re-sealed.	<b>Borings</b> : B-1 through B-35	Soil: TPH	None	2	No further action.
TF2-001	Aerial photograph 1951; Open burning is evident at Tank 21.	This AOC was GPS located and it was described as having minor, stressed vegetation. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-001-1 through TF2-001-5	<b>Soil</b> : Petroflag™ screening	None	1	Additional surface and subsurface soil sample collection and analysis for PAHs, dioxin and furans and metals.
TF2-002	Aerial photograph 1951; Large linear ground scar at	This AOC was GPS located and it was described as having stressed soils. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-001-2 through TF2-002-5	<b>Soil</b> : Petroflag™ screening	None	2	No further action.
TF2-003	Aerial photograph 1954; Open burning and stained ground at Tank 19.	This AOC was GPS located and it was described as a grassy, open area. A test pit was excavated and soil samples were screened for TPH using Petroflag™ and also sent to the laboratory for analysis. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-003-1 through TF2-003-5. Sent samples TF2-003-2, -3, -4 and -5 to laboratory for analysis.	<b>Soil:</b> DRO, Petroflag <sup>™</sup> screening	None	1	Additional surface and subsurface soil sample collection and analysis for PAHs, dioxin and furans and metals.
TF2-004	Aerial photograph 1954; Open burning and ground staining at Tank 19.	This AOC was GPS located and it was described as a grassy, open area. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-004-1 through TF2-004-7	<b>Soil:</b> Petroflag™ screening	None	1	Additional surface and subsurface soil sample collection and analysis for PAHs, dioxin and furans and metals.

# SUMMARY OF SITE CONDITIONS SAMPLING AND ANALYSIS PLAN REMEDIAL INVESTIGATION TANK FARM 2 NAVAL STATION, NEWPORT, NEWPORT, RHODE ISLAND

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Area	Description	History	Associated sampling locations	Available Data	Exceedances	Remedial Category	Other Information	Next Steps
TF2-005	Aerial photograph 1954; Open burning and ground staining at Tank 19.	This AOC was GPS located and it was described as a grassy, open area with a roadway partially within the AOC. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	TtEC 2005 soil evaloration: TE2-005-1	<b>Soil</b> : Petroflag™ screening	None	1		Additional surface and subsurface soil sample collection and analysis for PAHs, dioxin and furans and metals.
TF2-006	Aerial photograph 1954; Significant ground staining at Tank 22.	This AOC was GPS located and it was described as a grassy, open area. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	TtEC 2005 soil exploration:TF2-006-1 through TF2-006-7	<b>Soil</b> : Petroflag™ screening	None	2		No further action.
TF2-007	Aerial photograph 1954; Significant ground staining at Tank 22.	This AOC was GPS located and it was described as having low vegetation and a grassy, open area. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	TtFC 2005 soil exploration:TF2-007-1	<b>Soil</b> : Petroflag™ screening	None	2		No further action.
TF2-008	Aerial photograph 1954; Significant ground staining at Tank 22.	This AOC was GPS located and it was described as having loose rock and shale with no surface soils. Area that appeared to be staining on the 1954 aerial photographs is bedrock outcrop. No excavation of this area was completed.	None	None	Not Applicable	Not Applicable		No further action.
TF2-009	Aerial photograph 1954; Significant ground staining at Tank 27.	This AOC was GPS located and it was described as having low vegetation and no staining or stressed vegetation. A test pit was excavated and soil samples were screened for TPH using Petroflag™ and also sent to the laboratory for analysis. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	TtEC 2005 soil exploration:TF2-009-1 through TF2-009-9. Sent sample TF2-009-1	<b>Soil</b> : DRO, Petroflag <sup>™</sup> screening	None	2		No further action.
TF2-010	Aerial photograph 1954; Significant ground staining at Tank 27.	This AOC was GPS located and it was described as having stressed, stained soils in the SE area and otherwise, partially grassy and open and partially wooded. A test pit was excavated and soil samples were screened for TPH using Petroflag™ and also sent to the laboratory for analysis. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-010-1 through TF2-010-13. Sent samples TF2-010-3, -6 and -12 to laboratory for analysis.	<b>Soil</b> : DRO, Petroflag™ screening	None	2		No further action.
TF2-011	Aerial photograph 1962; Medium-toned material or liquid east of Tank 19.	This AOC was GPS located. The area was observed to be a fire hydrant and fire lines buried beneath a berm. The decision was made that no excavation was required. Aerial photographs showed anomalies here which turned out to be the buried fire protection infrastructure.	None	None	Not Applicable	Not Applicable		No further action.

# SUMMARY OF SITE CONDITIONS SAMPLING AND ANALYSIS PLAN REMEDIAL INVESTIGATION TANK FARM 2 NAVAL STATION, NEWPORT, NEWPORT, RHODE ISLAND

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Area	Description	History	Associated sampling locations	Available Data	Exceedances	Remedial Category	Other Information	Next Steps
TF2-012	Aerial photograph 1962:	This AOC was GPS located and it was described as an open area with minor stressed vegetation. A test pit was excavated and soil samples were screened for TPH using Petroflag™ and also sent to the laboratory for analysis. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	through TF2-012-5. Sent samples TF2-012-1	<b>Soil</b> : DRO, VOC, SVOC, Petroflag™ screening	None	2		No further action.
TF2-013	Aerial photograph 1962; Ground staining at Tank 28.	This AOC was GPS located and it was described as having no stressed vegetation or ground staining. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-013-1 through TF2-013-5	<b>Soil</b> : Petroflag™ screening	None	2		No further action.
TF2-014	Aerial photograph 1962; Ground staining at Tank 28.	This AOC was GPS located and it was described as an open vegetated area with no stress or staining. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-014-1 through TF2-014-5	<b>Soil</b> : Petroflag™ screening	None	2		No further action.
TF2-015	Aerial photograph 1962; Ground staining at Tank 29.	This AOC was GPS located and it was described as an open area that surrounds the Tank 29 blockhouse. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-015-1 through TF2-015-5	<b>Soil</b> : Petroflag™ screening	None	2		No further action.
TF2-016	Aerial photograph 1962; Staining and dark toned	This AOC was GPS located and it was described as not having any staining or mounded material that was observable. A test pit was excavated and soil samples were screened for TPH using Petroflag™ and also sent to the laboratory for analysis. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-016-1 through TF2-016-5. Sent samples TF2-016-3 and -4 to laboratory for analysis.	<b>Soil</b> : DRO, Petroflag <sup>™</sup> screening	None	2		No further action.
TF2-017	Aerial photograph 1962; Two pits are southwest of Tank 26.	This AOC was GPS located and it was described as having stressed vegetation and two, small craters and various small depressions. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.		<b>Soil</b> : Petroflag™ screening	None	2		No further action.

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Area	Description	History	Associated sampling locations	Available Data	Exceedances	Remedial Category	Other Information	Next Steps
TF2-018	Access road leads to two	This AOC was GPS located and it was described as having stressed vegetation. It was recommended to extend the test pit outside the area boundary. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-018-1 through TF2-018-5	<b>Soil</b> : Petroflag™ screening	None	2		No further action.
TF2-019	IAarial nhotograph 1964	This AOC was GPS located and it was described as having a depression in the middle of the AOC. It was stated that the test pit would be extended outside the area boundary. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-019-1 through TF2-019-5	<b>Soil:</b> Petroflag™ screening	None	2		No further action.
TF2-020	Ground scar is visible in	This AOC was GPS located. It was stated that the excavation would occur in AOC 20 and not AOC 39, which overlaps AOC 20. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	through TF2-020-5	<b>Soil</b> : Petroflag™ screening	None	2		No further action.
TF2-021	Aerial photograph 1972; Ground staining evident near Tank 19.	AOC was GPS located. Area that appeared to be staining in the 1972 aerial photographs is a bedrock outcrop. No excavation of this area was completed.	None	None	Not Applicable	Not Applicable		No further action.
TF2-022	access road west of Tanks 27 and 28. Containers remain in	defined by a 1'-1' concrete berm. A test pit was excavated	TtEC 2005 soil exploration:TF2-022-TP1-1 through TF2-022-TP1-5 and TF2-022-TP2-1 through TF2-022-TP2-5. Sent all samples to laboratory for analysis except TF2-022-TP2-2.	<b>Soil</b> : DRO, GRO, Petroflag <sup>™</sup> screening	None	2	JP-5 soils	No further action.
TF2-023	scarred or cleared areas are	This AOC was GPS located and it was described as having debris along the western edge. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-023-1 through TF2-023-5	<b>Soil</b> : Petroflag™ screening	None	2		No further action.

# SUMMARY OF SITE CONDITIONS SAMPLING AND ANALYSIS PLAN REMEDIAL INVESTIGATION TANK FARM 2 NAVAL STATION, NEWPORT, NEWPORT, RHODE ISLAND

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Area	Description	History	Associated sampling locations	Available Data	Exceedances	Remedial Category	Other Information	Next Steps
TF2-024	scarred or cleared areas are	This AOC was GPS located and it was described as being previously cleared but currently with heavy, new growth. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	through 1F2-024-9	<b>Soil</b> : Petroflag™ screening	None	2		No further action.
TF2-025	Ground scar and/or liquid are	GPS located area. Area that appeared to be ground staining/liquid in the 1975 aerial photograph is a bedrock outcrop. No excavation of this area was completed.	None	None	Not Applicable	Not Applicable		No further action.
TF2-026	probable liquid located north	heavy scrub vegetation present. A test pit was excavated and soil samples were screened for TPH using Petroflag™ and also sent to the laboratory for analysis. Test pit sample locations were surveyed with a GPS, the area was photographed and	TtEC 2005 soil exploration:TF2-026-1 through TF2-026-8. Also including TF2-026-	Groundwater: DRO, GRO, PAH, TPH, Lead, VOC Soil: DRO, GRO, VOC, SVOC, Petroflag™ screening	None	2	JP-5 soils	No further action.
TF2-027	Aerial photograph 1979; Hose leads from structure to a discharge point at Tank 23/Hose observed in aerial photos lead from T23 vault.	This AOC was GPS located and it was described as having distressed vegetation. A test pit was excavated and soil samples were screened for TPH using Petroflag™. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	TtEC 2005 soil exploration:TF2-027-1 through TF2-027-5	<b>Soil</b> : Petroflag™ screening	None	2		No further action.
TF2-028	Aerial photograph 1979; Large stained areas west of hose and discharge point.	· · ·	TtEC 2005 Remediation: TF2-028-9-1 through TF2-028-9-4. Sent samples TF2-028-	<b>Soil</b> : DRO, Petroflag <sup>™</sup> screening	No exceedences post-remedial excavation.	2		No further action.
TF2-029	Aerial photograph 1979; Large	excavated and soil samples were screened for TPH using	TtFC 2005 soil exploration:TF2-029-1	<b>Soil</b> : Petroflag™ screening	None	2		No further action.

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Area	Description	History	Associated sampling locations	Available Data	Exceedances	Remedial Category	Other Information	Next Steps
TF2-030	Aerial photograph 1981; Hose or pipe leads from a structure or pit at Tank 20.	This AOC was GPS located and it was described as having a depression on the western side of the Tank 23 blower. A test pit was excavated and soil samples were screened for TPH using Petroflag™ and also sent to the laboratory for analysis. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-030-1 through TF2-030-5. Sent sample TF2-030-4	<b>Soil</b> : DRO, VOC, SVOC, Petroflag™ screening	None	2		No further action.
TF2-031	Aerial photograph 1981; Hose or pipe leads from a structure or pit at Tank 20.	This AOC was GPS located and it was described as having minor stressed vegetation and a new, 4" hose on the ground. A test pit was excavated and soil samples were screened for TPH using Petroflag™ and also sent to the laboratory for analysis. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-031-1 through TF2-031-5. Sent all samples from test pit to laboratory for analysis.	<b>Soil</b> : DRO, Petroflag™ screening	None	2		No further action.
TF2-032	tank farm. Debris in area in	AOC was located with a GPS. The AOC was described as a fill area with large debris and slopes too steep to excavate. Fill made up of piles, tires, concrete. AOC investigated with the TF2-037 test pit.	See TF2-037	See TF2-037	See TF2-037	2		No further action.
TF2-033	Mound of dark toned material	GPS located area, mound present and will be excavated through the middle. Combined with AOCs 034 and 026 and investigated with the 026 test pit.	See TF2-026	See TF2-026	See TF2-026	2	JP-5 soils	No further action.
TF2-034	of staining or liquid adjacent	GPS located area. Area contains construction/ demolition debris including drum with stained soil. Combined with AOCs 033 and 026 and investigated with the 026 test pit.	See TF2-026	See TF2-026	See TF2-026	2	JP-5 soils	No further action.
TF2-035	Aerial photograph 1988; Light- toned mounded material is visible in the former container storage area.	This AOC was GPS located and the mound was present. It was stated that the test pit would run through the center of the mounded material. A test pit was excavated and soil samples were screened for TPH using Petroflag™ and also sent to the laboratory for analysis. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	<b>TtEC 2005 soil exploration</b> :TF2-035-1 through TF2-035-9. Sent samples TF2-035-1, 2, 5,6, and 9 to laboratory for analysis.	<b>Soil</b> : DRO, GRO, Petroflag™ screening	None	2	JP-5 soils	No further action.
TF2-036	toned mounded material is visible in the former container	This AOC was GPS located and the mound was present. It was stated that the test pit would run through the center of the mounded material. Combined with AOC-035, investigated with TF2-035 test pit.	See TF2-035	See TF2-035	See TF2-035	2	JP-5 soils	No further action.

# SUMMARY OF SITE CONDITIONS SAMPLING AND ANALYSIS PLAN REMEDIAL INVESTIGATION TANK FARM 2

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Area	Description	History	Associated sampling locations	Available Data	Fxceedances	Remedial Category	Other Information	Next Steps
TF2-037	Aerial photograph 1988; Two large pits (one containing a cylindrical object) are in the north central portion of TF2.	AOC was GPS located and described as an area on top of fill with stressed vegetation present. The test pit was excavated and samples collected and screened for TPH by Petroflag™ and also sent to the laboratory for analysis. Remedial excavations were performed east, west, and north of TF2-037-2. Confirmatory samples were collected and screened for TPH by Petroflag™ and also sent to the laboratory for analysis. All sample locations were surveyed with a GPS, the area was photographed and the test pits and excavation area was backfilled with clean material.	TtEC 2005 Remediation: TF2-037-2-1	<b>Soil</b> : DRO, VOC, SVOC, Petroflag™ screening	No exceedences post-remedial excavation.	2		No further action.
TF2-038	Aerial photograph 1988; Two large pits (one containing a cylindrical object) are in the north central portion of TF2.	AOC was located with a GPS. Area identified as a pit on the 1988 aerial photograph is a water valve. Will not excavate in this area.	None	None	Not Applicable	Not Applicable		No further action.
TF2-039	Iground scarred area are	AOC was located with a GPS. Area identified as three pits (TF2 039, TF2-040, TF2-041) on photographs are water valves, scarred area observed in photos due to installation of these valves. Will not excavate. AOC TF2-020, which overlaps TF2-039, was excavated.	See TF2-020	See TF2-020	See TF2-020	2		No further action.
TF2-040	Aerial photograph 1988; Three pits (one polygon) and a ground scarred area are within and adjacent to wooded area on the western side of TF2.	See TF2-039.	See TF2-039	See TF2-039	See TF2-039	2		No further action.
TF2-041	Aerial photograph 1988; Three pits (one polygon) and a ground scarred area are within and adjacent to wooded area on the western side of TF2.	See TF2-039.	See TF2-039	See TF2-039	See TF2-039	2		No further action.
TF2-042	Aerial photograph 1988; Three pits and a ground scarred area are within and adjacent to wooded area on the western side of TF2.	This AOC was GPS located and described as a ground scarred area and that the AOC follows drainage west, construction debris and water valve present. A test pit was excavated and soil samples were screened for TPH using Petroflag™ and also sent to the laboratory for analysis. Test pit sample locations were surveyed with a GPS, the area was photographed and the test pit was backfilled with the excavated material.	through TF2-042-9. Sent samples TF2-042-3	<b>Soil</b> : DRO, Petroflag™ screening	None	2		No further action.

# SUMMARY OF SITE CONDITIONS SAMPLING AND ANALYSIS PLAN REMEDIAL INVESTIGATION TANK FARM 2 NAVAL STATION, NEWPORT, NEWPORT, RHODE ISLAND PAGE 10 of 10

Area	Description	History	Associated sampling locations	Available Data	lExceedances	Remedial Category	Other Information Next Steps
TF2-043	Aerial photograph 1992; A large linear ground scar is on the west side of T26/Line from transformer building 219, vegetated area. Appears to be utility related as an underground line marker is present.	Appears to be utility related as an underground line marker	None	None	Not Applicable	Not Applicable	No further action.
Storage/ JP-5	Located west of Tank 28. Was previously investigaated as AOCs -022, -026, -033, -034, -035 and -036.	From historical photographs and documents it appears that this area was formerly used to store Naval Buoys and to store a pile of JP-5 impacted soil	see AOCs -022, -026, -033, -034, -035 and - 036	Soil and Groundwater: DRO/GRO, VOCs, SVOCs Groundwater: lead Soil: Petroflag <sup>TM</sup>	Not Applicable	1	Soil not tested for lead. Additional surface and subsurface soil sample collection and analysis for lead
Bldg 219	Former Electrical Service/ Transformer Building	This building was used for electrical equipment, including transformers.	SIRAR 2006: TF2-B219-1 through TF2-B219-4 surface soil samples collected around building perimeter all sent for laboratory analysis	soil: VOC, PCB	PCBs exceeded	1	Additional surface and subsurface soil sample collection and analysis for PCBs.

#### Notes

1. Tables A3.1 through A.3.14 of this Appendix include the specific criteria used to determine exceedances in the column titled 'Exceedances'.

# TABLE A-2 SUMMARY OF HISTORICAL GROUNDWATER SAMPLING SAMPLING AND ANALYSIS PLAN REMEDIAL INVESTIGATION TANK FARM 2 NAVAL STATION, NEWPORT, NEWPORT, RHODE ISLAND PAGE 1 of 2

Date	Activity	Wells	Analyses	Comments
October - December, 1996	GZA installed 11 monitoring wells	GZ-201 to GZ-211		
September and October, 1997	GZA installed an additional 17 monitoring wells	GZ-212 to GZ-228		
June, 1997	GZA collected groundwater samples.	GZ-201 through GZ-211	TPH, VOCs and PAHs	A water sample could not be collected from GZ-207 because it was dry.
December, 1997	GZA collected groundwater samples	GZ-212 through GZ-228 and GZ-207	TPH, VOCs and PAHs	
June, 1999	FwEC collected groundwater samples from nine of the 27 monitoring wells on site	GZ-207, GZ-208, GZ -212, GZ -213, GZ -214, GZ -218, GZ - 221, GZ -225 and GZ -227		
	Tatra Tash CC callected water	GZ-201 to GZ-208; GZ-210 to GZ -228 and RW-1 to RW-5.	VOCs, SVOCs and TPH DRO	
March - May 2005	Tetra Tech EC collected water samples from all "serviceable" monitoring wells, totalling 32 monitoring wells.	GZ-202 and GZ-208	Samples from these two wells were also analyzed as an oilwater mixture and the lab performed petroleum fingerprint analysis on the samples.	The samples were identified in the Number 2 Fuel Oil range.

# TABLE A-2 SUMMARY OF HISTORICAL GROUNDWATER SAMPLING SAMPLING AND ANALYSIS PLAN REMEDIAL INVESTIGATION TANK FARM 2 NAVAL STATION, NEWPORT, NEWPORT, RHODE ISLAND PAGE 2 of 2

Date	Activity	Wells	Analyses	Comments
July, 2005	In July 2005, Tetra Tech EC submitted a Work Plan for additional monitoring well installations at the Site. The plan proposed 20 additional monitoring wells, to be installed, if necessary, in the backfilled areas surrounding each tank. RIDEM provided comments on this Work Plan, but the work was never initiated, because it was determined that, based upon the results of the SIRAR, additional groundwater monitoring wells were not required.			
2009	Tetra Tech EC collected groundwater samples from 15 site wells. All groundwater	GZ-203,-205, -209, -210, -211, -212,-213, -215, -217, -218, - 219, -220, -222, -223, -225	VOCs, SVOCs, and total and dissolved lead	
2009	samples collected were analyzed for VOCs, SVOCs and total and dissolved lead.	Five selected wells: GZ-205, - 215, -219, -223, -225	TPH (GRO and DRO)	

#### Summary of Conditions at Tank 19 Tank Farm 2 NAVSTA, Newport, Rhode Island

#### **Groundwater**

			Sample ID	GZ-	-201	GZ-225		GZ-226
	<b>Collection Date</b>	03/11/05*	Jun-97	04/01/05	Jun-99	03/11/05		
Analyte	Unit	RIDEM GB GW Objectives	RIDEM GB GW Upper Concentration Limit					
1-Methylnaphthalene	ug/L	NS	NS	ND	10	ND	ND	ND
Bis(2-Ethylhexyl)Phthalate	ug/L	NS	NS	ND	ND	ND	6.0	ND
Diesel Range Organics (DRO)	mg/L	NS	NS	15	NT	0.15	NT	ND
Diethylphthalate	ug/L	NS	NS	ND	ND	ND	2.0J	ND
Di-N-Butylphthalate	ug/L	NS	NS	ND	ND	ND	1.0J	ND
Methylene Chloride	ug/L	NS	NS	ND	ND	ND	3.0J	ND
TPH	mg/L	NS	NS	NT	1600	NT	0.73	NT
VOCs	mg/L	NS	NS	ND	ND	ND	NA	ND

<sup>\*</sup>GZ-201 was analyzed at 10x dilution for VOCs

J=Estimated Value

NS= No Standard

NT = Not tested

NA = Not applicable

# **LNAPL** Gauging

	07.004	07.005	07.000
	GZ-201	GZ-225	GZ-226
Date	(ft)	(ft)	(ft)
Apr-09	NM	NG	ND
Feb-02	ND	ND	ND
Aug-01	ND	ND	ND
Jul-01	ND	ND	ND
Jun-99	ND	ND	NG
Mar-98	ND	NG	ND
Jul-97	0.12	NG	NG
Jun-97	0.12	NG	NG
May-97	ND	NG	NG
Apr-97	0.02	NG	NG
Mar-97	0.02	NG	NG
Feb-97	0.01	NG	NG
Jan-97	0.01	NG	NG

ND = Not detected

NG = Not gauged

NM = not measurable

## Soil

				Sample ID	GZ-201	GZ-225	GZ-226	B-35	TF2-Tankpit19-3
	Colle	ection Date	Nov-96	Oct-97	Oct-97	May-97	6/1/05		
Analyte	Unit	RIDEM Method 1 Industrial / Commercial Direct Exposure Criteria	Method 1 GB TPH Leachability Criteria	RIDEM Upper Concentr ation Limit	(0-2')	(0-2')	(4-6')	(10-12')	(10')
Bis(2-ethylhexyl)phthalate	mg/kg	410	NS	10,000	NT	NT	NT	NT	0.25J
DRO	mg/kg	2,500	NS	30,000	NT	NT	NT	NT	ND
PAHs	mg/kg	NS	NS	NS	ND	ND	ND	NT	ND
TPH	mg/kg	2,500	NS	30,000	ND	ND	40	ND	NT
VOCs	mg/kg	NS	NS	NS	ND	ND	ND	NT	ND

ND = Not detected above laboratory reporting limits

NS= No Standard

NT = Not tested

J=Estimated Value

ND = Not detected above laboratory reporting limits

Summary of Conditions at Tank 20 Tank Farm 2 NAVSTA, Newport, Rhode Island

#### Groundwater

			Sample ID	GZ-	202		GZ-218		RW-4
		Colle	ection Date	4/7/05	Jun-97	04/01/05*	Jun-99	Dec-97	
		RIDEM GB	RIDEM GB GW						
		GW	Upper						
		Objectives	Concentr						
Analyte	Unit		ation						
1,2,4-Trimethylbenzene	ug/L	NS	Limit NS	NT	ND	ND	7	5.3	
1-Methylnaphthalene	ug/L	NS	NS	NT	13	ND	ND	ND	
Bis(2-ethylHexyl)Phthalate	ug/L	NS	NS	NT	ND	ND	9	ND	
Diesel Range Organics (DRO)	mg/l	NS	NS	NT	NT	13	NT	NT	Not
Diethylphthalate	ug/L	NS	NS	NT	ND	ND	2.0J	ND	Sampled
Fluorene	ug/l	NS	NS	NT	ND	4	ND	ND	
Fuel Oil #2 (C9-C25)	mg/L	NS	NS	790**	NT	NT	NT	NT	
Methylene Chloride	ug/L	NS	NS	NT	ND	ND	3.0J	ND	
Naphthalene (VOC)	ug/L	NS	NS	NT	ND	ND	8	9.7	
PAHs	mg/L	NS	NS	NT	NA	NA	NA	ND	
Phenanthrene	ug/l	NS	NS	NT	ND	8.1	ND	ND	
P-Isopropyltoluene	ug/L	NS	NS	NT	ND	ND	ND	1.1	
TPH	mg/L	NS	NS	NT	2.1	NT	3.7	5.4	
VOCs	mg/L	NS	NS	NT	ND	ND	NA	NA	

<sup>\*</sup>GZ-218 was analyzed at 100x dilution for VOCs

ND = Not detected above laboratory reporting limits

J=Estimated Value

NS= No Standard

NT = Not tested

NA = Not applicable

# Soil

				Sample ID	GZ-202	GZ-218	B-8
			Colle	ection Date	Nov-96	Oct-97	May-97
Analyte	Unit	RIDEM Method 1 Industrial/C ommercial Direct Exposure Criteria	GBIPH	RIDEM Upper Concentra tion Limit	(0-2')	(5-7')	(10-12')
PAHs	mg/kg	NS	NS	NS	ND	ND	NT
TPH	mg/kg	2,500	2,500	30,000	ND	ND	ND
VOCs	mg/kg	NS	NS	NS	ND	ND	NT

ND = Not detected above laboratory reporting limits

NS= No Standard

NT = Not tested

# **LNAPL Gauging**

LIVAL L Gauging									
	GZ-202	GZ-218							
Date	(ft)	(ft)							
7/27/2009	0.35	NG							
7/9/2009	0.5	NG							
6/12/2009	1.13	NG							
6/1/2009	1.17	NG							
5/21/2009	0.79	NG							
5/6/2009	0.25	NG							
4/29/2009	0.4	NG							
4/13/2009	1.66	NG							
Spring 2005	ND	NG							
Feb-02	0.05	ND							
Nov-01	0.01	ND							
Oct-01	0.02	NG							
8/20/2001	0.05	NG							
Aug-01	NG	ND							
Jul-01	0.03	ND							
2001	0.05	NG							
Jun-99	0.28	ND							
Mar-98	0.23	NG							
Feb-98	ND	NG							
Jan-98	ND	NG							
Dec-97	0.02	ND							
Nov-97	0.03	NG							
Oct-97	0.11	NG							
Sep-97	0.02	NG							
Geb-at	0.02	110							
	ND	NG							
Aug-97									
Jul-97	ND	NG							
	detected	NG							
Jun-97									
May-97	ND	NG							
Apr-97	ND	NG							
Mar-97	ND	NG							
Feb-97	ND	NG							
Jan-97	ND	NG							
ND - Not doto	etod								

ND = Not detected

<sup>\*\*</sup>GZ-202 was sampled as oil-water mixture

#### Summary of Conditions at Tank 21 Tank Farm 2 NAVSTA, Newport, Rhode Island

#### Groundwater

	Sample ID						G	Z-227	RW-2
			<b>Collection Date</b>	5/19/2009	04/05/05	Jun-97	04/05/05*	Jun-99	4/14/2005
Analyte	Unit	RIDEM GB GW Objectives	RIDEM GB GW Upper Concentration Limit						
2-Methylnaphthalene	ug/L	NS	NS	ND	ND	ND	3.5	3.0J	ND
Bis(2-EthylHexyl)Phthalate	ug/L	NS	NS	ND	ND	ND	ND	4.0J	ND
Diesel Range Organics (DRO)	mg/L	NS	NS	NT	2.8	NT	3.1	NT	0.35
Lead (Dissolved)	ug/L	NS	NS	3.2 B	NT	NT	NT	NT	NT
Lead (Total)	ug/L	NS	NS	1.9 B	NT	NT	NT	NT	NT
Naphthalene (VOC)	ug/L	NS	NS	ND	ND	ND	ND	1.0J	ND
PAHs	mg/L	NS	NS	ND	ND	ND	NA	NA	ND
Phenanthrene	ug/L	NS	NS	ND	ND	ND	2.9	1.0J	ND
TPH	mg/L	NS	NS	NT	NT	ND	NT	1.6	NT
VOCs	mg/L	NS	NS	ND	ND	ND	ND	ND	ND

<sup>\*</sup>GZ227 was analyzed at 10x dilution for VOCs

ND = Not detected above laboratory reporting limits

J=Estimated Value

NS= No Standard

NT = Not tested

NA = Not applicable

## **LNAPL Gauging**

LIVAI L Gauging											
	GZ-203	RW-2	GZ-227								
Date	(ft)	(ft)	(ft)								
Apr-09	NG	NG	ND								
Feb-02	ND	ND	ND								
Nov-01	ND	ND	0.01								
Oct-01	NG	ND	ND								
Aug-01	ND	NG	0.02								
Jul-01	ND	NG	0.02								
2001	NG	NG	0.02								
Jun-99	ND	NG	ND								
Aug-97	ND	NG	NG								
Jul-97	ND	NG	NG								
Jun-97	ND	NG	NG								
May-97	ND	NG	NG								
Apr-97	ND	NG	NG								
Mar-97	ND	NG	NG								
Feb-97	ND	NG	NG								
Jan-97	ND	NG	NG								

ND = Not detected NG = Not gauged

## Soil

				Sample ID	GZ-203	GZ-200	GZ-227	B-31	TF2-Tank21-1	TF2-Tankpit21-1
	Collection Date						Oct-97	May-97	5/17/05	5/31/05
Analyte	Unit	RIDEM Method 1 Industrial / Commercial Direct Exposure Criteria	Method 1 GB TPH Leachability Criteria	RIDEM Upper Concentr ation Limit	Not Sampled	(0-2')	(0-2')	(10-12')	(1')	(5')
Diesel Range Organics (DRO)	mg/kg	2,500	NS	30,000		NT	NT	NT	48	120
PAHs	mg/kg	NS	NS	NS		ND	ND	NT	NT	NT
SVOCs	mg/kg	NS	NS	NS		ND	ND	NT	NT	ND
TPH	mg/kg	2,500	NS	30,000		11	ND	ND	NT	NT
VOCs	mg/kg	NS	NS	NS		ND	ND	NT	NT	ND

ND = Not detected above laboratory reporting limits

NS= No Standard

NT = Not tested

GZ-200 is a duplicate of GZ-227

#### Summary of Conditions at Tank 22 Tank Farm 2 NAVSTA, Newport, Rhode Island

#### Groundwater

		S	Sample ID	GZ-204	GZ-217	RW-5
		Collec	Jun-97	Dec-97		
Analyte	Unit	RIDEM GB GW Objectives	RIDEM GB GW Upper Concentration Limit			Not Sampled
PAHs	mg/L	NS	NS	ND	ND	
TPH	mg/L	NS	NS	ND	2	
VOCs	mg/L	NS	NS	ND	ND	

ND = Not detected above laboratory reporting limits

NS= No Standard

#### Soil

			Sa	mple ID	GZ-204	GZ-217	B-27	B-28
			Collecti	ion Date	Nov-96	Oct-97	5/1/97	5/1/97
Analyte	Unit	RIDEM Method 1 Industrial/Commercial Direct Exposure Criteria	Method 1 GB TPH Leachability Criteria	RIDEM Upper Concentration Limit	(5-6')	(0-2')	(10-12')	(10-12')
PAHs	mg/kg	NS	NS	NS	ND	ND	NT	NT
TPH	mg/kg	2,500	2,500	30,000	29	ND	ND	ND
VOCs	mg/kg	NS	NS	NS	ND	ND	NT	NT

ND = Not detected above laboratory reporting limits

NS= No Standard

NT = Not tested

# **LNAPL Gauging**

	GZ-204	GZ-217	RW-5
Date	(ft)	(ft)	(ft)
2/1/2002	ND	ND	ND
Nov-01	ND	ND	ND
Oct-01	NG	NG	ND
Aug-01	ND	ND	NG
Jul-01	ND	ND	NG
Jun-99	ND	ND	NG
Aug-97	ND	NG	NG
Jul-97	ND	NG	NG
Jun-97	ND	NG	NG
May-97	ND	NG	NG
Apr-97	ND	NG	NG
Mar-97	ND	NG	NG
Feb-97	ND	NG	NG
Jan-97	ND	NG	NG

ND = Not detected

#### Summary of Conditions at Tank 23 Tank Farm 2 NAVSTA, Newport, Rhode Island

#### Groundwater

			Sample ID		GZ-205		GZ-216			
		Colle	ction Date	5/20/09	3/11/05*	Jun-97	3/9/05	Dec-97		
Analyte	Unit	RIDEM GB GW Objective s	RIDEM GB GW Upper Concentr ation Limit							
1,3,5-Trimethylbenzene	ug/l	NS	NS	ND	ND	1.2	ND	ND		
1-Methylnaphthalene	ug/l	NS	NS	ND	ND	84	ND	ND		
2-Methylnaphthalene	ug/l	NS	NS	ND	35	10	ND	ND		
Acenaphthene	ug/l	NS	NS	ND	3.6	ND	ND	ND		
Diesel Range Organics (DRO)	mg/l	NS	NS	0.16	13	NT	0.35	NT		
Fluorene	ug/l	NS	NS	ND	6.0	ND	ND	ND		
Gasoline Range Organics (GRO)	mg/l	NS	NS	0.01	NT	NT	NT	NT		
Isopropylbenzene	ug/l	NS	NS	ND	ND	2.3	ND	ND		
Lead (Dissolved)	ug/l	NS	NS	ND	NT	NT	NT	NT		
Lead (Total)	ug/l	NS	NS	1.7B	NT	NT	NT	NT		
Naphthalene (SVOC)	ug/l	NS	NS	ND	ND	42	ND	ND		
Naphthalene (VOC)	ug/l	NS	NS	ND	6.4	37	ND	ND		
n-propylbenzene	ug/l	NS	NS	ND	ND	3.5	ND	ND		
o-xylene	ug/l	NS	NS	ND	ND	1.1	ND	ND		
PAHs	mg/l	NS	NS	ND	ND	NA	ND	ND		
Phenanthrene	ug/l	NS	NS	ND	11	ND	ND	ND		
sec-butylbenzene	ug/l	NS	NS	ND	ND	1.2	ND	ND		
TPH	mg/l	NS	NS	NT	NT	ND	ND	ND		
VOCs	mg/l	NS	NS	ND	ND	NA	ND	ND		

<sup>\*</sup>GZ-205 was analyzed at 50x dilution for VOCs

NS= No Standard

NT = Not tested

NA = Not applicable

# Soil

	Sample I													
	Collection Date													
Analyte	Unit	RIDEM Method 1 Industrial/Commercial Direct Exposure Criteria	Method 1 GB TPH Leachability Criteria	RIDEM Upper Concentration Limit	(25-27')		(10-12')							
1-Methylnaphthalene	mg/kg	NS	NS	10,000	2.9	Not Sampled	NT							
2-Methylnaphthalene	mg/kg	10,000	NS	10,000	1.5	Sampleu	NT							
Acenaphthene	mg/kg	10,000	NS	10,000	0.34		NT							
Fluorene	mg/kg	10,000	NS	10,000	0.86		NT							
Naphthalene (SVOC)	mg/kg	10,000	NS	10,000	0.41		NT							
naphthalene (VOC)	mg/kg	10,000	NS	10,000	1.4		NT							
n-butylbenzene	mg/kg	NS	NS	10,000	0.42		NT							
Phenanthrene	mg/kg	10,000	NS	10,000	2.5		NT							
sec-butylbenzene	mg/kg	NS	NS	10,000	0.3		NT							
TPH	mg/kg	2,500	2,500	30,000	930		ND							

ND = Not detected above laboratory reporting limits

NS= No Standard

NT = Not tested

# **LNAPL Gauging**

		ugiii
	GZ-205	GZ-216
Date	(ft)	(ft)
Dec-02	ND	NG
Feb-02	NG	ND
Aug-01	ND	ND
Jul-01	ND	ND
Jun-99	ND	ND
Aug-97	ND	NG
Jul-97	ND	NG
Jun-97	ND	NG
May-97	ND	NG
Apr-97	ND	NG
Mar-97	ND	NG
Feb-97	ND	NG
Jan-97	ND	NG

ND = Not detected

ND = Not detected above laboratory reporting limits

B = The result reported is less than reporting limit, but greater than instrument detection

Summary of Conditions at Tank 24 Tank Farm 2 NAVSTA, Newport, Rhode Island

#### Groundwater

	Sample ID GZ-206 GZ-220														
	Collection Date														
			RIDEM												
		RIDEM GB	GB GW												
		GW	Upper												
		Objectives	Concentr												
		Objectives	ation												
Analyte	Unit		Limit												
PAHs	mg/l	NS	NS	ND	ND										
sec-butylbenzene	ug/l	NS	NS	2.7	ND										
TPH	mg/l	NS	NS	ND	ND										
VOCs	mg/l	NS	NS	NA	ND										

ND = Not detected above laboratory reporting limits

NS= No Standard

NA = Not applicable

#### Soil

				Sample ID	GZ-206	GZ-220	B-14	B-13	TF2-Tank 24-2	TF2-Tank 24-2D
			Colle	ction Date	Nov-96	Oct-97	May-97	May-97	5/17/05	unknown
Analyte	Unit	RIDEM Method 1 Industrial/ Commerci al Direct Exposure Criteria	Method 1 GB TPH Leachabil ity Criteria	RIDEM Upper Concentr ation Limit	(10-12')	(8-8.7')	(10-12')	(10-12')	(1')	unknown
DRO	mg/kg	2,500	2,500	30,000	NT	NT	NT	NT	34	80
PAHs	mg/kg	NS	NS	NS	ND	ND	NT	NT	NT	NT
TPH	mg/kg	2,500	2,500	30,000	ND	ND	ND	ND	NT	NT
VOCs	mg/kg	NS	NS	NS	ND	ND	NT	NT	NT	NT

ND = Not detected above laboratory reporting limits

NS= No Standard

NT = Not tested

## **LNAPL Gauging**

	<u> </u>	··· <u> </u>
	GZ-206	GZ-220
Date	(ft)	(ft)
2/1/2002	ND	ND
Aug-01	ND	ND
Jul-01	ND	ND
Jun-99	NG	ND
Aug-97	ND	NG
Jul-97	ND	NG
35582	ND	NG
May-97	ND	NG
Apr-97	ND	NG
Mar-97	ND	NG
Feb-97	ND	NG
Jan-97	ND	NG
ND - Not dete	ected NG = No	t dalided

ND = Not detected NG = Not gauged

# Table A-3.7 Summary of Conditions at Tank 25 Tank Farm 2 NAVSTA, Newport, Rhode Island

#### Groundwater

			Sample ID	GZ-20	7		GZ-221			GZ-223		GZ-224
			<b>Collection Date</b>	4/1/05	Jun-99	4/13/05	Jun-99	Dec-97	5/20/09	4/1/05	Dec-97	4/1/05
Analyte	Unit	RIDEM GB GW Objectives	RIDEM GB GW Upper Concentration Limit									
Acetone	ug/l	NS	NS	ND	9.0J	ND	ND	ND	ND	ND	ND	ND
Bis(2-Ethylhexyl)Phthalate	ug/l	NS	NS	ND	4.0J	ND	2.0J	ND	ND	ND	ND	ND
Chloroform	ug/l	NS	NS	ND	ND	ND	ND	ND	ND	ND	1.3	ND
Diesel Range Organics (DRO)	mg/l	NS	NS	0.79	NT	4.9	NT	NT	0.12	0.23	NT	0.16
Diethylphthalate	ug/l	NS	NS	ND	1.0J	ND	ND	ND	ND	ND	ND	ND
Di-N-Butylphthalate	ug/l	NS	NS	ND	1.0J	ND	ND	ND	ND	ND	ND	ND
Isopropylbenzene	ug/l	NS	NS	1.3	ND	ND	ND	ND	ND	ND	ND	ND
Lead (Dissolved)	ug/l	NS	NS	NT	NT	NT	NT	NT	1.6B	NT	NT	NT
Lead (Total)	ug/l	NS	NS	NT	NT	NT	NT	NT	2.5B	NT	NT	NT
m,p-xylenes	ug/l	NS	NS	0.95	ND	ND	ND	ND	ND	ND	ND	ND
Methylene Chloride	ug/l	NS	NS	ND	4.0J	ND	ND	ND	ND	ND	ND	ND
Naphthalene (VOC)	ug/l	NS	NS	ND	3.0J	ND	ND	ND	ND	ND	ND	ND
n-butylbenzene	ug/l	NS	NS	0.58	ND	ND	ND	ND	ND	ND	ND	ND
n-propylbenzene	ug/l	NS	NS	1.4	ND	ND	ND	ND	ND	ND	ND	ND
o-xylenes	ug/l	NS	NS	2.4	ND	ND	ND	ND	ND	ND	ND	ND
PAHs	mg/l	NS	NS	ND	NA	ND	ND	ND	ND	ND	ND	ND
TPH	mg/l	NS	NS	NT	3.5	NT	0.7	ND	NT	NT	ND	NT
VOCs	mg/l	NS	NS	NA	NA	ND	ND	ND	ND	ND	NA	ND

ND = Not detected above laboratory reporting limits

J = Estimated Value

B = The result reported is less than reporting limit, but greater than instrument detection

NS= No Standard

NT = Not tested

NA = Not applicable

**LNAPL Gauging** 

	GZ-207	GZ-221	GZ-223	GZ-224
Date	(ft)	(ft)	(ft)	(ft)
Feb-02	ND	NG	ND	ND
Aug-01	ND	NG	ND	ND
Jul-01	ND	NG	ND	ND
Jun-05	NG	NG	NG	NG
Jun-99	ND	ND	ND	ND
Aug-97	ND	NG	NG	NG
Jul-97	ND	NG	NG	NG
Jun-97	ND	NG	NG	NG
May-97	ND	NG	NG	NG
Apr-97	ND	NG	NG	NG
Mar-97	ND	NG	NG	NG
Feb-97	ND	NG	NG	NG
Jan-97	ND	NG	NG	NG

ND = Not detected

NG = Not gauged

## Soil

Soil																																				
				Sample ID	GZ-221	GZ-223	GZ-224	B-10	B-12	TF2- Tank25-2	TF2-T25- R2	TF2-T25- R3	TF2-T25- R4	TF2-T25- R5	TF2-T25- R6	TF2-T25- R7	TF2-T25- R8	TF2-T25- R9	TF2-T25- R10B	TF2-T25- R11	TF2-T25- R12	TF2-T25- R14	TF2-T25- R16	TF2-T25- R18	TF2-T25- R19	TF2-T25- R20	TF2-T25- R23	TF2-T25- R27	TF2-T25- R30	TF2-T25- R32	TF2-T25- R33	TF2-T25- R34	TF2-T25- R35	TF2-T25- R36	TF2-T25- R37	TF2-T25- R38
				Collection Date	Oct-97	Oct-97	Oct-97	May-97	May-97	5/17/05									2/2/06								1/31/06	1/31/06	1/31/06	2/8/06	2/8/06	6/19/06	6/19/06	6/19/06	6/19/06	6/19/06
Analyte		RIDEM Method 1 Industrial / Commercial Direct Exposure Criteria	Method 1 GB TPH Leachability Criteria	RIDEM Upper Concentration Limit				,			sidewall of remedial excavation	sidewall of remedial excavation	base of remedial excavation	sidewall of remedial excavation	remedial	unknown	base of remedial excavation	sidewall of remedial excavation	base of remedial excavation	base(13') of remedial excavation	base of remedial excavation	base of remedial	sidewall of remedial excavation													
1,2,4-Trimethylbenzene	mg/kg	NS	NS	10,000	ND	ND	ND	NT	NT	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
1,3,5-Trimethylbenzene	mg/kg	NS	NS	10,000	ND	ND	ND	NT	NT	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
1-Methylnaphthalene	mg/kg	NS	NS	10,000	ND	ND	ND	NT	NT	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
2-Methylnaphthalene	mg/kg	10,000	NS	10,000	ND	ND	ND	NT	NT	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Anthracene	mg/kg	10000	NS	10,000	ND	ND	ND	NT	NT	0.14	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Benzo(a)anthracene	mg/kg	7.8	NS	10,000	ND	ND	ND	NT	NT	0.48	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Benzo(a)pyrene	mg/kg	0.8	NS	10,000	ND	ND	ND	NT	NT	0.48	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Benzo(b)fluoranthene	mg/kg	7.8	NS	10,000	ND	ND	ND	NT	NT	0.38	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Benzo(ghi)perylene	mg/kg	10000	NS	10,000	ND	ND	ND	NT	NT	0.31	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Benzo(k)fluoranthene	mg/kg	78	NS	10,000	ND	ND	ND	NT	NT	0.36	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Chrysene	mg/kg	780	NS	10,000	ND	ND	ND	NT	NT	0.5	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
DRO (Diesel Range Organics)		2,500	2,500	30,000	NT	NT	NT	NT	NT	210	71	140	9.9	450	17	210	12	44	13	9	380	350	110	430	82	340	11	6	20	ND	8.1	200	97	17	300	77
Fluoranthene	mg/kg	10000	NS	10,000	ND	ND	ND	NT	NT	1.0	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Fluorene	mg/kg	10,000	NS	10,000	ND	ND	ND	NT	NT	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Indeno(1,2,3-cd)pyrene	mg/kg	7.8	NS	10,000	ND	ND	ND	NT	NT	0.25	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Naphthalene (SVOC)	mg/kg	10,000	NS	10,000	ND	ND	ND	NT	NT	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Naphthalene (VOC)	mg/kg	10,000	NS	10,000	ND	ND	ND	NT	NT	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
PAHs	mg/kg	NS	NS	NS	ND	ND	ND	NT	NT	NA	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Phenanthrene	mg/kg	10,000	NS	10,000	ND	ND	ND	NT	NT	0.55	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
Pyrene	mg/kg	10000	NS	10,000	ND	ND	ND	NT	NT	1.2	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND
SVOC	mg/kg	NS	NS	NS	ND	ND	ND	NT	NT	NA	NT	NT	NT	NT	NT	NT	NT	NT	ND	NT	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND						
TPH	mg/kg	2,500	2,500	30,000	ND	ND	200	1,700	550	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT	NT
VOCs	mg/kg	NS	NS	NS	ND	ND	ND	NT	NT	ND	NT	NT	NT	NT	NT	NT	NT	NT	ND	NT	ND	ND	ND	ND	ND	ND	ND	ND	ND	ND						

ND = Not detected above laboratory reporting limits

NS= No Standard

NT = Not tested NA = Not applicable

Only currently applicable soil data is shown. Any samples from areas removed during the remedial excavation were not included in this table.

#### Summary of Conditions at Tank 26 Tank Farm 2 NAVSTA, Newport, Rhode Island

#### Groundwater

			Sample ID	(	3Z-208		GZ-	RW-3	
		Co	lection Date		Jun-99	Jun-97	5/20/09		
Analyte	Unit	RIDEM GB GW Objectives	RIDEM GB GW Upper Concentrat ion Limit						
1,2,4-Trimethylbenzene	ug/L	NS	NS	NT	16	ND	ND	ND	
1-Methylnaphthalene	ug/L	NS	NS	NT	ND	43	ND	ND	
2-Methylnaphthalene	ug/L	NS	NS	NT	6	ND	ND	ND	
bis(2-ethylhexyl)phthalate	ug/L	NS	NS	NT	4.0J	ND	ND	ND	
Fluorene	ug/L	NS	NS	NT	2.0J	ND	ND	ND	Not
Fuel Oil #2 (C9-C25)	mg/L	NS	NS	740**	NT	NT	NT	NT	Sampled
Lead (Dissolved)	ug/L	NS	NS	NT	NT	NT	ND	NT	
Lead (Total)	ug/L	NS	NS	NT	NT	NT	1.5 B	NT	
Naphthalene (SVOC)	ug/L	NS	NS	NT	8	ND	ND	ND	
Naphthalene (VOC)	ug/L	NS	NS	NT	20	ND	ND	ND	
PAHs	mg/L	NS	NS	NT	ND	NA	ND	ND	
Phenanthrene	ug/L	NS	NS	NT	3.0J	ND	ND	ND	
TPH	mg/L	NS	NS	NT	37	2	NT	0.36	
VOCs	mg/L	NS	NS	NT	NA	ND	ND	ND	

<sup>\*\*</sup>GZ-208 was sampled as oil-water mixture.

ND = Not detected above laboratory reporting limits

J = Estimated Value

B = The result reported is less than reporting limit, but greater than instrument detection

NS= No Standard

NT = Not tested

NA = Not applicable

## Soil

				Sample ID	GZ-208	GZ-222	B-16					
	Collection Date											
Analyte	RIDEM Method 1 Industrial/ ommercia Direct Exposure Unit Criteria		Method 1 GB Leachabilit y Criteria	RIDEM Upper Concentra tion Limit	(5-7')	(5-7')	(10-12')					
PAHs	mg/kg	NS	NS	NS	ND	ND	NT					
TPH	mg/kg	2,500	2,500	30,000	ND	ND	ND					
VOCs	mg/kg	NS	NS	NS	ND	ND	NT					

ND = Not detected above laboratory reporting limits

NS= No Standard

NT = Not tested

**LNAPL** Gauging

LNAPL	Gaugi									
	GZ-208	GZ-222								
Date	(ft)	(ft)								
7/27/2009	1.25	NG								
7/9/2009	0.31	NG								
6/12/2009	0.25	NG								
6/1/2009	0.1	NG								
5/21/2009	0.13	NG								
5/6/2009	0.05	NG								
4/29/2009	0.31	NG								
4/13/2009	0.42	NG								
Spring 2005	ND	NG								
Feb-02	0.06	ND								
Nov-01	0.41	ND								
Oct-01	0.03	NG								
8/28/2001	0.02	NG								
8/20/2001	0.08	NG								
Aug-01	NG	ND								
Jul-01	0.1	ND								
2001	0.01	NG								
Jun-99	ND	ND								
Mar-98	ND	NG								
Feb-98	ND	NG								
Jan-98	ND	NG								
Dec-97	0.01	NG								
Nov-97	ND	NG								
Oct-97	0.01	NG								
Sep-97	0.01	NG								
Aug-97	0.02	NG								
Jul-97	ND	NG								
Jun-97	detected	NG								
May-97	ND	NG								
Apr-97	ND	NG								
Mar-97	ND	NG								
Feb-97	ND	NG								
Jan-97	ND	NG								
ND = Not detect	ed	<u> </u>								

ND = Not detected

#### Summary of Conditions at Tank 27 Tank Farm 2 NAVSTA, Newport, Rhode Island

#### Groundwater

			Sample ID	GZ-209		GZ-2	12	GZ-	-219
			<b>Collection Date</b>	Jun-97	5/21/09	Jun-99	Dec-97	5/20/09	Dec-97
			RIDEM GB GW						
		RIDEM GB	Upper						
		GW	Concentration						
Analyte	Unit	Objectives	Limit						
Diesel Range Organics (DRO)	mg/l	NS	NS	NT	NT	NT	NT	0.06	NT
1,2,4-Trimethylbenzene	ug/l NS		NS	ND	ND	ND	1.6	ND	ND
1-Methylnaphthalene	ug/l	NS	NS	33	ND	ND	ND	ND	ND
2-Methylnaphthalene	ug/l	NS	NS	10	ND	ND	ND	ND	ND
Bis(2-EthylHexyl)Phthalate	ug/l	NS	NS	ND	ND	2.0 JB	ND	ND	ND
Chloroform	ug/l	NS	NS	ND	ND	ND	1.1	ND	ND
Lead (Dissolved)	ug/l	NS	NS	NT	ND	NT	ND	2.1B	NT
Lead (Total)	ug/l NS		NS	NT	ND NT		ND	1.6B	NT
PAHs			NS	ND	ND	ND	ND	ND	ND
TPH	mg/l NS		NS	2.5	NT	0.45	ND	NT	ND
VOCs	mg/l	NS	ND	ND	ND	NA	ND	ND	

ND = Not detected above laboratory reporting limits

J = Estimated Value

B = The result reported is less than reporting limit, but greater than instrument detection

NS= No Standard

NT = Not tested

NA = Not applicable

## Soil

Sample ID					GZ-209	B-18	TF2-Tank27-1	TF2-Tankpit27-2	TF2-Tankpit27-5
Collection Date					Oct-96	May-97	5/17/05	5/31/05	5/31/05
Analyte (Soil)	Unit	RIDEM Method 1 Industrial/ Commercia I Direct Exposure Criteria	Method 1 GB Leachability Criteria	RIDEM Upper Concent ration Limit	(15-17')	(10-12')	(1')	(5')	(5')
1,2,3-Trichlorobenzene	mg/kg		NS	10,000	0.103	NT	NT	NT	ND
1,2,4-Trimethylbenzene	mg/kg	NS	NS	10,000	0.0441	NT	NT	NT	ND
1,3,5-Trimethylbenzene	mg/kg	NS	NS	10,000	0.01	NT	NT	NT	ND
1-Methylnaphthalene	mg/kg	NS	NS	10,000	20	NT	NT	NT	ND
2-Methylnaphthalene	mg/kg		NS	10,000		NT	NT	NT	ND
Diesel Range Organics (DRO)	mg/kg	2,500	2,500	30,000	NT	NT	41	ND	9.4
Fluorene	mg/kg	10,000	NS	10,000	5.6	NT	NT	NT	ND
Naphthalene (SVOC)	mg/kg	10,000	NS	10,000	4.9	NT	NT	NT	ND
Naphthalene (VOC)	mg/kg	10,000	NS	10,000	0.031	NT	NT	NT	ND
Phenanthrene	mg/kg	10,000	NS	10,000	9.6	NT	NT	NT	ND
TPH	mg/kg	2,500	500 2,500 30,000			ND	NT	NT	ND

ND = Not detected above laboratory reporting limits

NS= No Standard

NT = Not tested

# **LNAPL** Gauging

GZ-209 GZ-212 GZ-219													
	GZ-209	GZ-212	GZ-219										
Date	(ft)	(ft)	(ft)										
Feb-02	ND	ND	ND										
Aug-01	ND	ND	ND										
Jul-01	ND	ND	ND										
Jun-99	ND	ND	ND										
Aug-97	ND	NG	NG										
Jul-97	ND	NG	NG										
Jun-97	ND	NG	NG										
May-97	ND	NG	NG										
Apr-97	ND	NG	NG										
Mar-97	ND	NG	NG										
Feb-97	ND	NG	NG										
Jan-97	ND	NG	NG										

ND = Not detected

#### Summary of Conditions at Tank 28 Tank Farm 2 NAVSTA, Newport, Rhode Island

#### Groundwater

Groundwater						
			Sample ID	GZ-210	GZ-	213
		Col	lection Date	Jun-97	Jun-99	Dec-97
Analyte	RIDEM GB		RIDEM GB GW Upper Concentrat ion Limit			
1,2,4-Trimethylbenzene	ug/l	NS	NS	22	ND	ND
1,3,5-Trimethylbenzene	ug/l	NS	NS	1.8	ND	ND
1-methylnaphthalene	ug/l	NS	NS	44	ND	ND
Benzene	ug/l	140	18,000	4.2	ND	ND
bis(2-ethylhexyl)phthalate	ug/l	NS	NS	NT	4.0JB	ND
Ethylbenzene	ug/l	1600	16,000	4.9	ND	ND
isopropylbenzene	ug/l	NS	NS	3.4	ND	ND
m+p xylene	ug/l	NS	NS	1.7	ND	ND
n-propylbenzene	ug/l	NS	NS	5.5	ND	ND
PAHs	mg/l	NS	NS	NT	ND	ND
p-isopropylbenzene	ug/l	NS	NS	6.4	ND	ND
TPH	mg/l	NS	NS	ND	0.3	1.1
VOCs	mg/l	NS	NS	NA	ND	ND

ND = Not detected above laboratory reporting limits

NS= No Standard

J=Estimated Value

B = The result reported is less than reporting limit, but greater than instrument detection

NT = Not tested

NA = Not applicable

#### Soil

	Sample ID													
	ction Date	Nov-96		May-97										
Analyte	Unit	RIDEM Method 1 Industrial/C ommercial Direct Exposure Criteria	Method 1 GB TPH Leachabilit y Criteria	RIDEM Upper Concentr ation Limit	(5-7')	Not Sampled	(10-12')							
PAHs	mg/kg	NS	NS	NS	ND		NT							
TPH	mg/kg	2,500	2,500	30,000	ND		18							
VOCs	mg/kg	NS	NS	NS	ND		NT							

ND = Not detected above laboratory reporting limits

NS= No Standard

NT = Not tested

**LNAPL** Gauging

-		
	GZ-210	GZ-213
Date	(ft)	(ft)
4/13/2009	ND	NG
Feb-02	ND	ND
Aug-01	ND	ND
Jul-01	ND	ND
Jun-99	ND	ND
Jun-97	ND	NG
May-97	ND	NG
Apr-97	ND	NG
Mar-97	ND	NG
Feb-97	ND	NG
Jan-97	ND	NG

ND = Not detected

#### Summary of Conditions at Tank 29 Tank Farm 2 NAVSTA, Newport, Rhode Island

#### Groundwater

			Sample ID		GZ-211		GZ-228	RW-1
		Colle	ection Date	5/20/2009	03/11/05*	Jun-97	3/9/2005	
Analyte	Unit	RIDEM GB GW Objective s	RIDEM GB GW Upper Concentr ation Limit	Not Sampled				
1,2,4-Trimethylbenzene	ug/L NS ug/L NS		NS	because product	47	ND	ND	
1-Methylnaphthalene			NS	was	ND	51	ND	Not Sampled
2-Methylnaphthalene	ug/L	NS	NS	observed	1500	12	ND	
Diesel Range Organics (DRO)	mg/L	NS	NS	in the	1200	NT	0.22	
Fluorene	ug/L	NS	NS	sample	500	20	ND	
Phenanthrene	ug/L	NS	NS	tubing	900	32	ND	
Pyrene	ug/L	NS	NS		120	ND	ND	
TPH	mg/L	NS	NS		NT	190	NT	
VOCs	mg/L	NS	NS		NA	ND	ND	

<sup>\*</sup>GZ211 was analyzed at 25x dilution for VOCs, and 20x for SVOC

NS= No Standard

NT = Not Tested

NA = Not Applicable

#### Soil

				Sample ID	GZ-211	GZ-228	B-25	TF2-Tank29-1
		Dec-96	Oct-97	May-97	05/17/05			
Analyte	Unit	RIDEM Method 1 Industrial /Commer cial Direct Exposure Criteria	Method 1 GB TPH Leachabil ity Criteria	RIDEM Upper Concentra tion Limit	(0-2')	(0-2')	(10-12')	(1')
Diesel Range Organics (DRO)	mg/kg	2,500	2,500	30,000	NT	NT	NT	26
PAHs	mg/kg	NS	NS	NS	ND	ND	NT	NT
TPH	mg/kg	2,500	2,500	30,000	ND	ND	ND	NT
VOCs	mg/kg NS		NS	NS	ND	ND	NT	NT

ND = Not detected above laboratory reporting limits

NS= No Standard

NT = Not Tested

# **LNAPL** Gauging

	Oddging	)
	GZ-211	GZ-228
Date	(ft)	(ft)
4/13/2009	product observed during sampling	NG
Feb-02	0.01	ND
Nov-01	0.46	ND
Oct-01	0.17	ND
Aug-01	NG	ND
Jul-01	NG	0.01
2001	0.11	NG
Jun-99	0.08	ND
Mar-98	ND	NG
Feb-98	0.01	NG
Jan-98	0.02	NG
Dec-97	0.03	NG
10/97-11/97	0.02	NG
Sep-97	0.01	NG
Aug-97	ND	NG
Jul-97	ND	NG
Jun-97	detected	NG
May-97	ND	NG
Apr-97	ND	NG
Mar-97	ND	NG
Feb-97	ND	NG
Jan-97	ND	NG
ND = Not detecte	ed	

ND = Not detected NG = Not Gauged

ND = Not detected above laboratory reporting limits

# Summary of Conditions along Fuel Transport Piping Tank Farm 2 NAVSTA, Newport, Rhode Island

# Soil

			Sample ID	B-1	B-2	B-3	B-4	B-5	B-6	B-7	B-8	B-9	B-10	B-11	B-12	B-13	B-14	B-15	B-16	B-17	B-18	B-19	B-20	B-21	B-22	B-23	B-24	B-25	B-26	B-27	B-28	B-29	B-30	B-31	B-32	B-33	B-34	B-35
		Colle	ection Date	May-97	May-97	May-97	May-97	7 May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97	May-97
Analyte	Unit	RIDEM Method 1 Industrial/ Commerc ial Direct Exposure Criteria	Upper	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12') (Excavated during the remediation of Tank 25)	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')	(10-12')
TPH	mg/kg	2,500 2,500	30,000	20	63	<10	<10	28	<10	<10	<10	<10	1700	1800	550	<10	<10	<10	<10	<10	<10	<10	<10	<10	18	<10	<10	<10	<10	<10	<10	<10	<10	<10	<10	<10	<10	<10

ND = Not detected above laboratory reporting limits

NS= No Standard

#### Summary of Conditions at Former JP-5 Soil Piles/ Buoy Storage Area Tank Farm 2 NAVSTA, Newport, Rhode Island

## Groundwater

Ol Galla Water			Sample ID		
	GZ-215				
	5/21/2009	Dec-97			
			RIDEM		
		DIDEM	GB GW		
	RIDEM U		Upper		
	GB GW				
		Objective	ation		
Analyte	Unit	S	Limit		
Diesel Range Organics (DRO)	mg/L	NS	NS	0.06	NT
Gasoline Range Organics (GRO)	mg/L	NS	NS	ND	NT
PAHs	mg/L	NS	NS	ND	ND
TPH	mg/L	NS	NS	NT	1.3
Lead (Dissolved)	ug/L	NS	NS	0.9B	NT
Lead (Total)	ug/L	NS	NS	2.1B	NT
p-isopropyltoluene	ug/L	NS	NS	ND	2.1
sec-butylbenzene	ug/L	NS	NS	ND	3.3

ND = Not detected above laboratory reporting limits

NS= No Standard

NT= Not Tested

# **LNAPL** Gauging

	<u> </u>
	GZ-215
Date	(ft)
4/13/2009	sheen
Feb-02	ND
Nov-01	ND
Oct-01	ND
Aug-01	ND
Jul-01	ND
Jun-99	ND
_	

ND = Not detected

# Soil

Sample ID					GZ-215	TF2-026-1	TF2-026-2A	TF2-026-3A	TF2-026-4	TF2-026-5	TF2-026-6	TF2-026-7	TF2-026-8A	TF2-035-1	TF2-035-2	TF2-035-5	TF2-035-6	TF2-035-9
Collection Date				5/23/05	7/5/05	7/5/05	5/23/05	5/23/05	5/23/05	5/23/05	7/5/05	5/24/05	5/24/05	5/24/05	5/24/05	5/24/05		
Analyte	Unit	RIDEM Method 1 Industrial /Commer cial Direct Exposure Criteria	Method 1 GB TPH Leachabil ity Criteria	RIDEM Upper Concentr ation Limit		(1')	(1')	(1')	(1')	(1')	(2')	(1')	(1')	(1')	unknown	(2')	(0.5')	(1')
2-Butanone (Methyl Ethyl Ketone)	mg/kg	10,000	NS	10,000		NT	ND	0.062	NT	ND	NT	ND	ND	NT	NT	NT	NT	NT
2-Methylnaphthalene	mg/kg	10,000	NS	10,000		NT	ND	0.13J	NT	ND	NT	ND	ND	NT	NT	NT	NT	NT
Acetone	mg/kg	10,000	NS	10,000	Not	NT	ND	0.4	NT	ND	NT	ND	ND	NT	NT	NT	NT	NT
Benzo(a)anthracene	mg/kg	7.8	NS	10,000	Sampled	NT	ND	0.2	NT	ND	NT	ND	ND	NT	NT	NT	NT	NT
Benzo(a)pyrene	mg/kg	0.8	NS	10,000		NT	ND	0.21	NT	ND	NT	ND	ND	NT	NT	NT	NT	NT
Benzo(b)fluoranthene	mg/kg	7.8	NS	10,000		NT	ND	0.26	NT	ND	NT	ND	ND	NT	NT	NT	NT	NT
Benzo(k)fluoranthene	mg/kg	78	NS	10,000		NT	ND	0.17	NT	ND	NT	ND	ND	NT	NT	NT	NT	NT
Chrysene	mg/kg	780	NS	10,000		NT	ND	0.27	NT	ND	NT	ND	ND	NT	NT	NT	NT	NT
Fluoranthene	mg/kg	10000	NS	10,000		NT	ND	0.27	NT	ND	NT	ND	ND	NT	NT	NT	NT	NT
Phenanthrene	mg/kg	10,000	NS	10,000		NT	ND	0.19	NT	ND	NT	ND	ND	NT	NT	NT	NT	NT
Pyrene	mg/kg	10000	NS	10,000		NT	ND	0.66	NT	ND	NT	ND	0.13J	NT	NT	NT	NT	NT
SVOCs	mg/kg	NS	NS	NS		NT	ND	NA	NT	ND	NT	ND	NA	NT	NT	NT	NT	NT
TPH (DRO+GRO)	mg/kg	2,500	2,500	30,000		15*	1100*	488	94	211	45*	203	746	16*	74	8.4*	38*	25*
VOCs	mg/kg	NS	NS	NS		NT	ND	NA	NT	ND	NT	ND	ND	NT	NT	NT	NT	NT

ND = Not detected above laboratory reporting limits

NS= No Standard

'\*' = Only DRO were detected.

NT = Not tested

NA = Not applicable

J=Estimated Value

B = The result reported is less than reporting limit, but greater than instrument detection

#### Summary of Conditions at B219 Tank Farm 2 NAVSTA Newport, RI

# Soil

		TF2-B219-1	TF2-B219-2	TF2-B219-3	TF2-B219-4			
			6/2/05	6/2/05	6/2/05	6/2/05		
Analyte	Unit	RIDEM Method 1 Residenti al Direct Exposure Criteria	RIDEM Method 1 Industrial/ Commerc ial Direct Exposure Criteria	_	(0-0.5')	(0-0.5')	(0-0.5')	(0-0.5')
Chlorinated Benzenes	mg/kg	NS	NS	NS	ND	ND	ND	ND
PCB (Aroclor 1260)	mg/kg	10	10	10	18	0.33	4.1	11

ND = Not detected above laboratory reporting limits

NS= No Standard

ND(0.210) = not detected at indicated laboratory reporting limit

# APPENDIX B TETRA TECH AND EPA SOPS



**TETRA TECH NUS, INC.** 

# **STANDARD OPERATING PROCEDURES**

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Effective Date 01/29/01	Revision 2

Applicability

Tetra Tech NUS, Inc.

Prepared

Management Information Systems Department

Approved

D. Senovich



Subject DATABASE RECORDS AND QUALITY ASSURANCE

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#### 1.0 PURPOSE

The purpose of this document is to specify a consistent procedure for the quality assurance review of electronic and hard copy databases. This SOP outlines the requirements for establishment of a Database Record File, Quality Assurance review procedures, and documentation of the Quality Assurance Review Process.

#### 2.0 SCOPE

The methods described in this Standard Operating Procedure (SOP) shall be used consistently for all projects managed by Tetra Tech NUS (TtNUS).

# 3.0 GLOSSARY

<u>Chain-of-Custody Form</u> - A Chain-of-Custody Form is a printed form that accompanies a sample or a group of samples from the time of sample collection to the laboratory. The Chain-of-Custody Form is retained with the samples during transfer of samples from one custodian to another. The Chain-of-Custody Form is a controlled document that becomes part of the permanent project file. Chain-of-Custody and field documentation requirements are addressed in SOP SA-6.1.

<u>Electronic Database</u> - A database provided on a compact laser disk (CD). Such electronic databases will generally be prepared using public domain software such as DBase, RBase, Oracle, Visual FoxPro, Microsoft Access, Paradox, etc.

<u>Hardcopy Database</u> - A printed copy of a database prepared using the software discussed under the definition of an electronic database.

Form I - A printed copy of the analytical results for each sample.

<u>Sample Tracking Summary</u> - A printed record of sample information including the date the samples were collected, the number of samples collected, the sample matrix, the laboratory to which the samples were shipped, the associated analytical requirements for the samples, the date the analytical data were received from the laboratory, and the date that validation of the sample data was completed.

# 4.0 RESPONSIBILITIES

<u>Database Records Custodian</u> - It shall be the responsibility of the Database Records Custodian to update and file the Sample Tracking Summaries for all active projects on a weekly basis. It shall be the responsibility of the Database Records Custodian to ensure that the most recent copies of the Sample Tracking Summaries are placed in the Database Records file. It shall be the responsibility of the Database Records Custodian to ensure that a copy of all validation deliverables is provided to the Project Manager (for placement in the project file). It shall be the responsibility of the Database Records Custodian to ensure that photocopies of all validation deliverables and historical data and reports (as applicable) are placed in the Database Records file.

<u>Data Validation Coordinator</u> - It shall be the responsibility of the Data Validation Coordinator (or designee) to ensure that the Sample Tracking Summaries are maintained by the Database Records Custodian. It shall be the responsibility of the Data Validation Coordinator (or designee) to ensure that photocopies of all data validation deliverables are placed in the applicable Database Records file by the Database Records Custodian.

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<u>Earth Sciences Department Manager</u> - It shall be the responsibility of the Earth Sciences Department Manager (or equivalent) to ensure that all field personnel are familiar with the requirements of this Standard Operating Procedure (specifically Section 5.5).

<u>FOL</u> - It shall be the responsibility of the FOL (FOL) of each project to ensure that all field technicians or sampling personnel are thoroughly familiar with this SOP, specifically regarding provision of the Chain-of-Custody Forms to the Database Records Custodian. Other responsibilities of the FOL are described in Sections 5.4 and 5.5.

Management Information Systems (MIS) Manager - It shall be the responsibility of the MIS Manager to ensure that copies of original electronic deliverables (CDs) are placed in both the project files and the Database Records File. It shall be the responsibility of the MIS Manager (or designee) to verify the completeness of the database (presence of all samples) in both electronic and hardcopy form in the Database Records File. It shall be the responsibility of the MIS Manager to ensure that Quality Assurance Reviews are completed and are attested to by Quality Assurance Reviewers. It shall be the responsibility of the MIS Manager to ensure that records of the Quality Assurance review process are placed in the Database Records File. It shall be the responsibility of the MIS Manager to ensure that both electronic and hardcopy forms of the final database are placed in both the project and the Database Record File. It shall be the responsibility of the MIS Manager to ensure that data validation qualifiers are entered in the database.

Furthermore, it shall be the responsibility of the MIS Manager to participate in project planning at the request of the Project Manager, specifically with respect to the generation of level of effort and schedule estimates. To support the project planning effort, the MIS Manager shall provide a copy of the MIS Request From included as Attachment A to the project manager. It shall be the responsibility of the MIS Manager to generate level of effort and budget estimates at the time database support is requested if a budget does not exist at the time of the request. The MIS Request Form shall be provided to the Project Manager at the time of any such requests. It shall be the responsibility of the MIS Manager to notify the Project Manager of any anticipated level of effort overruns or schedule noncompliances as soon as such problems arise along with full justification for any deviations from the budget estimates (provided they were generated by the MIS Manager). It shall be the responsibility of the MIS Manager to document any changes to the scope of work dictated by the Project Manager, along with an estimate of the impact of the change on the level of effort and the schedule.

<u>Program/Department Managers</u> - It shall be the responsibility of the Department and/or Program Managers (or designees) to inform their respective department's Project Managers of the existence and requirements of this SOP.

Project Manager - It shall be the responsibility of each Project Manager to determine the applicability of this SOP based on: (1) program-specific requirements, and (2) project size and objectives. It shall be the responsibility of the Project Manager (or designee) to ensure that the FOL is familiar with the requirements regarding Chain-of-Custody Form provision to the Database Records Custodian. It shall be the responsibility of the Project Manager (or designee) to determine which, if any, historical data are relevant and to ensure that such data (including all relevant information such as originating entity, sample locations, sampling dates, etc.) are provided to the Database Records Custodian for inclusion in the Database Records File. It shall be the responsibility of the Project Manager to obtain project planning input regarding the level of effort and schedule from the MIS Manager. It shall be the responsibility of the Project Manager to complete the database checklist (Attachment A) to support the level of effort and schedule estimate and to facilitate database preparation and subroutine execution.

<u>Risk Assessment Department Manager</u> - It shall be the responsibility of the Risk Assessment Department Manager to monitor compliance with this Standard Operating Procedure, to modify this SOP as necessary, and to take corrective action if necessary. Monitoring of the process shall be completed on a quarterly basis.

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Quality Assurance Reviewers - It shall be the responsibility of the Quality Assurance Reviewers to verify the completeness of the sample results via review of the Chain-of-Custody Forms and Sample Tracking Summaries. It shall be the responsibility of the Quality Assurance Reviewers to ensure the correctness of the database via direct comparison of the hardcopy printout of the database and the hardcopy summaries of the original analytical data (e.g., Form Is provided in data validation deliverables). Correctness includes the presence of all relevant sample information (all sample information fields), agreement of the laboratory and database analytical results, and the presence of data validation qualifiers.

<u>Quality Manager</u> - It shall be the responsibility of the Quality Manager to monitor compliance with this Standard Operating Procedure via routine audits.

#### 5.0 PROCEDURES

#### 5.1 Introduction

Verification of the accuracy and completeness of an electronic database can only be accomplished via comparison of a hardcopy of the database with hardcopy of all relevant sample information. The primary purposes of this SOP are to ensure that 1) all necessary hardcopy information is readily available to Quality Assurance Reviewers; 2) ensure that the Quality Assurance review is completed in a consistent and comprehensive manner, and; 3) ensure that documentation of the Quality Assurance review process is maintained in the project file.

# 5.2 <u>File Establishment</u>

A Database Record file shall be established for a specific project at the discretion of the Project Manager. Initiation of the filing procedure will commence upon receipt of the first set of Chain-of-Custody documents from a FOL or sampling technician. The Database Record Custodian shall establish a project-specific file for placement in the Database Record File. Each file in the Database Record File shall consist of standard components placed in the file as the project progresses. Each file shall be clearly labeled with the project number, which shall be placed on the front of the file drawer and on each and every hanging file folder relevant to the project. The following constitute the minimum components of a completed file:

- Electronic Deliverables
- Sample Tracking Forms
- Chain-of-Custody Forms
- Data Validation Letters
- Quality Assurance Records

# 5.3 Electronic Deliverables

The format of electronic deliverables shall be specified in the laboratory procurement specification and shall be provided by the laboratory. The integrity of all original electronic data deliverables shall be maintained. This shall be accomplished via the generation of copies of each electronic deliverable provided by the laboratory. The original electronic deliverable shall be provided to the project manager for inclusion in the project file. A copy of the original electronic deliverable shall be placed in the Database Record File. The second copy shall be maintained by the MIS Manager (or designee) to be used as a working copy.

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# 5.4 Sample Tracking Forms

Updated versions of the sample tracking form for each relevant project shall be maintained by the Database Record Custodian. The Sample Tracking Forms shall be updated any time additional Chain-of-Custody Forms are received from a FOL or sampling technician, or at any time that data are received from a laboratory, or at any time that validation of a given data package (sample delivery group) is completed. The Data Validation Coordinator shall inform the Database Record Custodian of the receipt of any data packages from the laboratory and of completion of validation of a given data package to facilitate updating of the Sample Tracking Form. The Database Record Custodian shall place a revised copy of the Sample Tracking Form in the Database Record File anytime it has been updated. Copies of the updated Sample Tracking Form shall also be provided to the project manager to apprise the project manager of sample package receipt, completion of validation, etc.

# 5.5 Chain-of-Custody Forms

The Chain-of-Custody Forms for all sampling efforts will be used as the basis for (1) updating the Sample Tracking Form, and (2) confirming that all required samples and associated analyses have been completed. It shall be the responsibility of the FOL (or sample technician) to provide a photocopy of all Chain-of-Custody Forms to the Database Record Custodian immediately upon completion of a sampling effort. The Database Record Custodian shall then place the copies of the Chain-of-Custody Form(s) in the Database Record File. Upon receipt of a sample data package from an analytical laboratory, the Data Validation Coordinator shall provide a copy of the laboratory Chain-of-Custody Form to the Database Record Custodian. The Database Record Custodian shall use this copy to update the Sample Tracking Summary and shall place the copy of the laboratory-provided Chain-of-Custody Form in the Database Record File. The photocopy of the laboratory-provided Chain-of Custody Form shall be stapled to the previously filed field copy. Upon receipt of all analytical data, two copies of the Chain-of-Custody will therefore be in the file. Review of the Chain-of-Custody Forms will therefore be a simple mechanism to determine if all data have been received. Chain-of-Custody is addressed in SOP SA-6.1.

# 5.6 <u>Data Validation Letters</u>

All data validation deliverables (or raw data summaries if validation is not conducted) shall be provided for inclusion in both the Database Record File and the project file. If USEPA regional- or client-specific requirements are such that Form Is (or similar analytical results) need not be provided with the validation deliverable, copies of such results must be appended to the deliverable. It is preferable, although not essential that the validation qualifiers be hand-written directly on the data summary forms. The data validation deliverables (and attendant analytical summaries) will provide the basis for direct comparison of the database printout and the raw data and qualifiers.

# 5.7 Historical Data

At the direction of the Project Manager, historical data may also be included in a project-specific analytical database. In the event that historical data are germane to the project, hardcopy of the historical data must be included in the Database Record File. Historical data may be maintained in the form of final reports or as raw data. The information contained in the historical data file must be sufficient to identify its origin, its collection date, the sample location, the matrix, and any and all other pertinent information. All available analytical data, Chain-of-Custody Forms, boring logs, well construction logs, sample location maps, shall be photocopied by the Project Manager (or designee) and placed in one or more 3-ring binders. All information shall be organized chronologically by matrix. It shall be the responsibility of the Project Manager (or designee) to ensure that all inconsistencies between analytical data, Chain-of-Custody Forms, boring logs, sample log sheets, and field logbooks are identified and corrected. The Project Manager (or designee) shall decide which nomenclature is appropriate and edit, initial and date all relevant forms. Data entry may only be performed on information that has undergone the aforementioned

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editing process, thereby having a direct correlation between hardcopy information and what will become the electronic database.

# 6.0 RECORDS

Records regarding database preparation and quality assurance review include all those identified in the previous section. Upon completion of the database task, records from the file will be forwarded to the Project Manager for inclusion in the project file, or will be placed in bankers boxes (or equivalent) for storage. The final records for storage shall include the following minimum information on placards placed on both the top and end of the storage box:

Database Record File
PROJECT NUMBER:
SITE NAME:
DATE FILED://
SUMMARY OF CONTENTS ENCLOSED
BOX OF

Project- or program-specific record keeping requirements shall take precedence over the record keeping requirements of this SOP.

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# **ATTACHMENT A**



# MIS REQUEST FORM

Tetra Tech NUS, Inc.		
Project Name:		Request Date:
сто:		Date Data Available for Production:
Project Manager:		Request in Support of:
Requestor:	-	Database Lead:
Program/Client:		GIS Lead:
State/EPA Region	n:	Statistics Lead:
		Risk Lead:
Site Name(s) (Are	ea, OU, etc.):	
Sampling Date(s)	:	
Matrix:	☐GW ☐SO ☐SD [	SW Other:
Labels:	Labels needed for an upcomin	
	ted Hours	Additional Instructions:
Due Da		·
	Complete ETS Charge No.	
	FOL	
		Y
Data Entry:	C Observing data and to be and	
	Chemical data needs to be ent	
	Chemical data needs to be for	
	Field analytical data needs to b	
	Geologic data needs to be entered.  Hydrology data needs to be en	
Estimo	ted Hours	Additional Instructions:
Due Da		Additional instructions.
	Complete ETS Charge No.	:
	Complete E13 Onlarge No.	
Tables:	Full Data Printout	,
Tables.	Summary of Positive Hits	
	Occurance and Distribution	with criteria
	Sampling Analytical Summary	
	Other:	<u> </u>
Estima	ted Hours	Additional Instructions:
Due Da		
	Complete ETS Charge No.	
		:
GIS:	General Facility Location	
	Site Location	
	Potentiometric Contours/Groun	ndwater Flow
	Sample Location Proposed	
	Sample Location Existing	
	Tag Map Single Round	
	Tag Map Multiple Round	
	Chart Map	:
	3D Visualization	!
	EGIS CD	
	Other:	
	ted Hours	Additional Instructions:
Due Da		_ <del></del>
	Complete ETS Charge No.	
	T-T-V	;
Statistics:	Yes	Additional facts of one
	ted Hours	Additional Instructions:
Due Da		
	Complete ETS Charge No.	:
Geostatistics:	l Yes	
	ted Hours	Additional Instructions:
Due Da		- Additional High deligits
Due Da	Complete ETS Charge No.	
	Complete L10 Onarge 110.	
	<del></del>	



**TETRA TECH NUS, INC.** 

# STANDARD OPERATING PROCEDURES

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ity Tetra Tech NUS, Inc.

Prepared

Earth Sciences Department

Approved

D. Senovich L

Subject

SITE RECONNAISSANCE

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# 1.0 PURPOSE

The purpose of a site reconnaissance is to collect both general and technical information which will support the scoping, scheduling, implementing project activities, and writing reports for an environmental investigation. This procedure is not intended as a guide for Phase I investigations or for Environmental Baseline Survey activities.

# 2.0 SCOPE

This procedure is applicable to the performance of a site reconnaissance for initial site characterization. The steps necessary to develop and carry out a site reconnaissance are presented here. These steps include a list of equipment and items which may be needed, areas of special interest during field observations, and methods by which the field observation team can ensure that necessary and appropriate observations have been made.

# 3.0 GLOSSARY

<u>Site reconnaissance</u>. An onsite inspection program used to identify site-specific conditions that control scheduling, manpower, and affect costs. A site reconnaissance usually consists of visual observations and, often, the use of field monitoring instruments to identify potential health and safety threats and potential sampling locations for site evaluation during subsequent field investigations.

# 4.0 RESPONSIBILITIES

<u>Field Operations Leader (FOL)</u> is responsible for ensuring that the survey is carried out in sufficient detail. To accomplish this, the FOL must assign the proper personnel and equipment to characterize the site adequately, in accordance with the requirements defined in this procedure and best engineering practices. Other disciplines which may be applicable include (but are not limited to): Geology/Hydrogeology; Health and Safety; Ecological Specialists; and/or Engineering. In addition, the FOL is responsible for supervising equipment preparation, including necessary calibrations, and supervising field data collection and documentation in accordance with the methods described in all referenced standard operation procedures.

Project Manager is responsible for the following:

- Supervising the retrieval and examination of available, applicable information regarding the site.
- Obtaining appropriate program approvals and ensuring the preparation of a site Health and Safety plan for the site reconnaissance.
- Coordinating the field activities with the client and regulatory agencies, as applicable.

<u>Field Personnel</u> are primarily responsible for observing and documenting, either through written documentation or photographic evidence, the site reconnaissance. Field personnel will take direction from the FOL.

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# 5.0 PROCEDURES

# 5.1 Equipment Items/Needed

Below is a list of items that may be useful when conducting a site reconnaissance. All, or a portion of these items may be required, depending upon the objective of the site reconnaissance.

- · Health and Safety equipment and information as required by the Site Safety Officer.
- Maps (U.S.G.S. quadrangle, geologic maps, street and highway maps, and client facility maps).
- Geologic tools (compass, tape measure, hand level, camera, etc.).
- Physical monitoring equipment, if applicable (PID, Immunoassay Test Kits, etc.)
- Regional publications (U.S.G.S reports, water well surveys, U.S.D.A. soil conservation surveys, etc.).
- Site-specific publications by previous investigators (EPA aerial photographic analyses, remedial investigation reports, data on waste disposal practices, boring logs, etc.).
- Marking items (ink markers, surveyor's flagging, spray paint, pin flags, wooden stakes).
- Field notebooks.
- Local telephone book with yellow pages (for obtaining utilities, site trailer, living accommodations, etc.).

Sufficient time will be required in order to obtain some of the aforementioned material. In general, most publications can be obtained in time to be used in the site reconnaissance if ordered approximately 2 weeks before the actual site visit takes place.

# 5.2 Observations

A site reconnaissance usually requires one to two days, however, additional time may be needed depending upon the objective, site size, etc. The following observations, when applicable, should be documented either on a site map, field notebook, or photographed.

- General Site Access. It should be noted whether site roads provide access to all proposed work
  locations, or if it will be necessary to prepare access roads with either a backhoe, dozer, chain saws,
  etc., in order to get drill rigs, excavators, or other work vehicles to specific locations. If temporary
  driveways must be constructed from existing public roads, regulatory permits may be required.
  Military facilities may have specific security requirements which require detailed clearance procedures.
- Location of the Command Post or Site Trailer and Sanitary Facilities. The ideal location for the site trailer and sanitary facilities is a level area, within an uncontaminated zone, and centralized in order to provide easy access to work areas on the site. However, certain utility companies may require that the site trailer be placed within a specified radius (usually 100 feet), of the nearest utility pole. Contact the necessary utility companies and inquire about the requirements regarding service before conducting the site reconnaissance. Information that may be required by the utility companies is: type of electric service needed (inquire with trailer vendor for this information); and utility pole number of interest (pole numbers are usually stamped on a brass plate on the pole).

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- <u>Potable Water Sources</u>. Local fire departments may allow access to fire hydrants. Private water delivery companies may also be available in the area.
- Sources of Possible Contamination. Drums, tanks, sludge areas, areas of stressed vegetation, fill
  areas, and leachate seeps may indicate where sources of contamination exist. Filler pipes protruding
  from the ground surface may indicate the presence of underground storage tanks. Areas where the
  original ground surface has been reworked may be contaminated fill areas that have since been
  buried and covered with natural material. Previous environmental investigations may also identify
  source areas.
- <u>Location of Decon Areas and Storage/Disposal Areas for Equipment and Wastes Generated by Field Activities.</u>
- Locations of Surface Water Bodies. The locations of surface water bodies, both man-made and natural, and their relation to topographic highs may give an indication of the groundwater flow direction in the area (groundwater flow typically follows topography with the topographic highs serving as groundwater recharge areas, and the surface waters at topographic lows serve as groundwater discharge areas). Visible signs of contamination, the existence of aquatic life, flow rates, and approximate levels should also be observed and noted. Check if the surface water bodies could potentially be impacted by field activities. If so, appropriate sedimentation and erosion controls will be required.
- Existing Wells. Existing monitoring wells, or domestic wells within the site and off site, should be noted on a map, and access checked to see if the wells can be used for data collection.
- Outcrops. Outcrops can be useful in providing hydrogeologic data (lithologic description, strike and dip information, fracture and joint system analysis, identification of moist zones, etc.) Outcrops may occur naturally or be a part of a man-made feature such as a road-cut.
- <u>Lineaments</u>. A lineament is a straight lengthy feature on the earth's surface which is expressed topographically as a line of depression. Stream beds, vegetation patterns or soil characteristics may be aligned or controlled by this feature. Lineaments are due in some cases to the presence of intense jointing or faults beneath the ground surface. Groundwater in the bedrock may follow lineaments. Lineaments should be noted on site maps and described in the field notebooks.
- Bench or Property Markers. Benchmarks or property markers should be marked with paint or surveyor's flagging if encountered during a site reconnaissance. Surveyors may need to use these markers as a reference point when surveying. Benchmarks are typically a brass plate secured in concrete in the ground with numbering on the top. Property markers can range from a stake driven into the ground to a rock protruding from the ground surface. Facility contacts may also be aware of local benchmarks used during the course of other environmental or public work projects.
- <u>Metal Cultural Effects</u>. Overhead power lines, railroad tracks, junk automobiles, fences, etc. will greatly affect certain geophysical surveys. These features should be noted while conducting a site reconnaissance.

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# 6.0 RECORDS

The data collected during a site reconnaissance may have to be compiled into a trip report when returning from the field. This trip report can then be distributed to the project team. A site reconnaissance checklist is located in Attachment A which can be copied and used while conducting the site reconnaissance.

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# **ATTACHMENT A**

#### SITE RECONNAISSANCE CHECKLIST

# SITE SKETCH

Include the following as appropriate:

- Site Name
- Site location
- Site Boundaries
- Entrance locations
- · Access Roads and Security Requirements
- Disposal locations
- Storage areas
- Office areas
- Well locations
- Treatment facility locations
- Surface drainage, outcrops, general topography descriptions
- Cultural interferences

# CHEMICAL STORAGE FACILITIES DESCRIPTION

- Storage tanks numbers, volumes, condition, contents, etc.
- Drums number, conditions, labeling, etc.
- Lagoons and surface pits number, size, use of liner, contents, etc.

# TREATMENT SYSTEMS

Note the presence of any treatment systems. These can be difficult to evaluate visually. One should appraise general appearance, maintenance and visual integrity; ask operators for any monitoring records; note presence of odors; and visually characterize any effluents or residues. Describe type of wastes and volumes treated.

- Incinerators
- Flocculation/filtration
- · Chemical/physical treatment
- Biological treatment
- Volume reduction
- Waste recycling
- Compositing
- Other

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# **DISPOSAL FACILITIES**

Note the presence and use of any of the following operations. Include a description of the size, use of liners, soil type, and the presence of leachate. Provide a description of management practices. Interview site workers if possible. Describe waste types.

- Landfills
- Land forms
- Open dump
- Surface impoundment
- Underground injection
- Incineration

Also, records for disposal of concentrated/containerized waste should be reviewed.

#### HAZARDOUS SUBSTANCE CHARACTERISTICS

Ask facility contacts for manifests, inventories, or monitoring reports. Note markings on containers.

- Chemical identities
- Quantities
- Hazard characteristics (toxic, explosive, flammable, etc.)
- Container markings
- Monitoring data, other analytical data
- Physical state (liquid, solid, gas, sludge)

# **CHEMICAL PROCESS INFORMATION**

- Manufacturing processes and chemicals
- Off-specification or by-product disposal processes
- Housekeeping practices
- Locations of Plant Operations

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# HYDROGEOLOGIC ASSESSMENT

Look for situations that promote hazardous substance migration, i.e., porous soils, fractured bedrock formations, shallow water table and karst features.

- Soil type
- Surface water features
- Surface drainage pattern
- Outcrop studies
- Water wells (use, water depth, and construction details)
- Erosion potential
- Flooding potential
- Climatology

# **IDENTIFICATION OF SENSITIVE RECEPTORS**

- · Number and locations of private homes
- · Public buildings including tenant usage
- Areas of dead or dying vegetation or animals
- Presence of sensitive ecosystems (wetlands, tidal marshes, etc.)
- Other public use areas (roads, parks, etc.)
- Natural areas



**TETRA TECH NUS, INC.** 

# STANDARD OPERATING PROCEDURES

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Applicability

Tetra Tech NUS, Inc.

Prepared

Earth Sciences Department

Approved

D. Senovich

Subject

BOREHOLE AND SAMPLE LOGGING

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# 1.0 PURPOSE

The purpose of this document is to establish standard procedures and technical guidance on borehole and sample logging.

# 2.0 SCOPE

These procedures provide descriptions of the standard techniques for borehole and sample logging. These techniques shall be used for each boring logged to provide consistent descriptions of subsurface lithology. While experience is the only method to develop confidence and accuracy in the description of soil and rock, the field geologist/engineer can do a good job of classification by careful, thoughtful observation and by being consistent throughout the classification procedure.

# 3.0 GLOSSARY

None.

#### 4.0 RESPONSIBILITIES

<u>Site Geologist</u>. Responsible for supervising all boring activities and assuring that each borehole is completely logged. If more than one rig is being used on site, the Site Geologist must make sure that each field geologist is properly trained in logging procedures. A brief review or training session may be necessary prior to the start up of the field program and/or upon completion of the first boring.

# 5.0 PROCEDURES

The classification of soil and rocks is one of the most important jobs of the field geologist/engineer. To maintain a consistent flow of information, it is imperative that the field geologist/engineer understand and accurately use the field classification system described in this SOP. This identification is based on visual examination and manual tests.

# 5.1 <u>Materials Needed</u>

When logging soil and rock samples, the geologist or engineer may be equipped with the following:

- Rock hammer
- Knife
- Camera
- Dilute hydrochloric acid (HCI)
- Ruler (marked in tenths and hundredths of feet)
- Hand Lens

# 5.2 Classification of Soils

All data shall be written directly on the boring log (Figure 1) or in a field notebook if more space is needed. Details on filling out the boring log are discussed in Section 5.5.

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#### 5.2.1 USCS Classification

Soils are to be classified according to the Unified Soil Classification System (USCS). This method of classification is detailed in Figure 1 (Continued).

This method of classification identifies soil types on the basis of grain size and cohesiveness.

Fine-grained soils, or fines, are smaller than the No. 200 sieve and are of two types: silt (M) and clay (C). Some classification systems define size ranges for these soil particles, but for field classification purposes, they are identified by their respective behaviors. Organic material (O) is a common component of soil but has no size range; it is recognized by its composition. The careful study of the USCS will aid in developing the competence and consistency necessary for the classification of soils.

Coarse-grained soils shall be divided into rock fragments, sand, or gravel. The terms sand and gravel not only refer to the size of the soil particles but also to their depositional history. To insure accuracy in description, the term rock fragments shall be used to indicate angular granular materials resulting from the breakup of rock. The sharp edges typically observed indicate little or no transport from their source area, and therefore the term provides additional information in reconstructing the depositional environment of the soils encountered. When the term "rock fragments" is used it shall be followed by a size designation such as " $(1/4 \text{ inch}\Phi-1/2 \text{ inch}\Phi)$ " or "coarse-sand size" either immediately after the entry or in the remarks column. The USCS classification would not be affected by this variation in terms.

#### 5.2.2 Color

Soil colors shall be described utilizing a single color descriptor preceded, when necessary, by a modifier to denote variations in shade or color mixtures. A soil could therefore be referred to as "gray" or "light gray" or "blue-gray." Since color can be utilized in correlating units between sampling locations, it is important for color descriptions to be consistent from one boring to another.

Colors must be described while the sample is still moist. Soil samples shall be broken or split vertically to describe colors. Samplers tend to smear the sample surface creating color variations between the sample interior and exterior.

The term "mottled" shall be used to indicate soils irregularly marked with spots of different colors. Mottling in soils usually indicates poor aeration and lack of good drainage.

Soil Color Charts shall not be used unless specified by the project manager.

# 5.2.3 Relative Density and Consistency

To classify the relative density and/or consistency of a soil, the geologist is to first identify the soil type. Granular soils contain predominantly sands and gravels. They are noncohesive (particles do not adhere well when compressed). Finer-grained soils (silts and clays) are cohesive (particles will adhere together when compressed).

The density of noncohesive, granular soils is classified according to standard penetration resistances obtained from split-barrel sampling performed according to the methods detailed in Standard Operating Procedures GH-1.3 and SA-1.3. Those designations are:

Designation	Standard Penetration Resistance (Blows per Foot)
Very loose	0 to 4
Loose	5 to 10
Medium dense	11 to 30
Dense	31 to 50
Very dense	Over 50

Standard penetration resistance is the number of blows required to drive a split-barrel sampler with a 2-inch outside diameter 12 inches into the material using a 140-pound hammer falling freely through 30 inches. The sampler is driven through an 18-inch sample interval, and the number of blows is recorded for each 6-inch increment. The density designation of granular soils is obtained by adding the number of blows required to penetrate the last 12 inches of each sample interval. It is important to note that if gravel or rock fragments are broken by the sampler or if rock fragments are lodged in the tip, the resulting blow count will be erroneously high, reflecting a higher density than actually exists. This shall be noted on the log and referenced to the sample number. Granular soils are given the USCS classifications GW, GP, GM, SW, SP, SM, GC, or SC (see Figure 1).

The consistency of cohesive soils is determined by performing field tests and identifying the consistency as shown in Figure 2.

Cohesive soils are given the USCS classifications ML, MH, CL, CH, OL, or OH (see Figure 1).

The consistency of cohesive soils is determined either by blow counts, a pocket penetrometer (values listed in the table as Unconfined Compressive Strength), or by hand by determining the resistance to penetration by the thumb. The pocket penetrometer and thumb determination methods are conducted on a selected sample of the soil, preferably the lowest 0.5 foot of the sample in the split-barrel sampler. The sample shall be broken in half and the thumb or penetrometer pushed into the end of the sample to determine the consistency. Do not determine consistency by attempting to penetrate a rock fragment. If the sample is decomposed rock, it is classified as a soft decomposed rock rather than a hard soil. Consistency shall not be determined solely by blow counts. One of the other methods shall be used in conjunction with it. The designations used to describe the consistency of cohesive soils are shown in Figure 2.

# 5.2.4 Weight Percentages

In nature, soils are comprised of particles of varying size and shape, and are combinations of the various grain types. The following terms are useful in the description of soil:

Terms of Identifying Proportion of the Component	Defining Range of Percentages by Weight
Trace	0 - 10 percent
Some	11 - 30 percent
Adjective form of the soil type (e.g., "sandy")	31 - 50 percent

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# FIGURE 2 CONSISTENCY FOR COHESIVE SOILS

Consistency	Standard Penetration Resistance (Blows per Foot)	Unconfined Compressive Strength (Tons/Sq. Foot by pocket penetration)	Field Identification
Very soft	0 to 2	Less than 0.25	Easily penetrated several inches by fist
Soft	2 to 4	0.25 to 0.50	Easily penetrated several inches by thumb
Medium stiff	4 to 8	0.50 to 1.0	Can be penetrated several inches by thumb with moderate effort
Stiff	8 to 15	1.0 to 2.0	Readily indented by thumb but penetrated only with great effort
Very stiff	15 to 30	2.0 to 4.0	Readily indented by thumbnail
Hard	Over 30	More than 4.0	Indented with difficulty by thumbnail

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# Examples:

- Silty fine sand: 50 to 69 percent fine sand, 31 to 50 percent silt.
- Medium to coarse sand, some silt: 70 to 80 percent medium to coarse sand, 11 to 30 percent silt.
- Fine sandy silt, trace clay: 50 to 68 percent silt, 31 to 49 percent fine sand, 1 to 10 percent clay.
- Clayey silt, some coarse sand: 70 to 89 percent clayey silt, 11 to 30 percent coarse sand.

#### 5.2.5 Moisture

Moisture content is estimated in the field according to four categories: dry, moist, wet, and saturated. In dry soil, there appears to be little or no water. Saturated samples obviously have all the water they can hold. Moist and wet classifications are somewhat subjective and often are determined by the individual's judgment. A suggested parameter for this would be calling a soil wet if rolling it in the hand or on a porous surface liberates water, i.e., dirties or muddies the surface. Whatever method is adopted for describing moisture, it is important that the method used by an individual remains consistent throughout an entire drilling job.

Laboratory tests for water content shall be performed if the natural water content is important.

# 5.2.6 Stratification

Stratification can only be determined after the sample barrel is opened. The stratification or bedding thickness for soil and rock is depending on grain size and composition. The classification to be used for stratification description is shown in Figure 3.

# 5.2.7 Texture/Fabric/Bedding

The texture/fabric/bedding of the soil shall be described. Texture is described as the relative angularity of the particles: rounded, subrounded, subangular, and angular. Fabric shall be noted as to whether the particles are flat or bulky and whether there is a particular relation between particles (i.e., all the flat particles are parallel or there is some cementation). The bedding or structure shall also be noted (e.g., stratified, lensed, nonstratified, heterogeneous varved).

# 5.2.8 Summary of Soil Classification

In summary, soils shall be classified in a similar manner by each geologist/engineer at a project site. The hierarchy of classification is as follows:

- Density and/or consistency
- Color
- Plasticity (Optional)
- Soil types
- Moisture content
- Stratification
- Texture, fabric, bedding
- Other distinguishing features

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FIGURE 3
BEDDING THICKNESS CLASSIFICATION

Thickness (metric)	Thickness (Approximate English Equivalent)	Classification
> 1.0 meter	> 3.3'	Massive
30 cm - 1 meter	1.0' - 3.3'	Thick Bedded
10 cm - 30 cm	4" - 1.0'	Medium Bedded
3 cm - 10 cm	1" - 4"	Thin Bedded
1 cm - 3 cm	2/5" - 1"	Very Thin Bedded
3 mm - 1 cm	1/8" - 2/5"	Laminated
1 mm - 3 mm	1/32" - 1/8"	Thinly Laminated
< 1 mm	<1/32"	Micro Laminated

(Weir, 1973 and Ingram, 1954)

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# 5.3 Classification of Rocks

Rocks are grouped into three main divisions: sedimentary, igneous and metamorphic. Sedimentary rocks are by far the predominant type exposed at the earth's surface. The following basic names are applied to the types of rocks found in sedimentary sequences:

- Sandstone Made up predominantly of granular materials ranging between 1/16 to 2 mm in diameter.
- Siltstone Made up of granular materials less than 1/16 to 1/256 mm in diameter. Fractures irregularly. Medium thick to thick bedded.
- Claystone Very fine-grained rock made up of clay and silt-size materials. Fractures irregularly. Very smooth to touch. Generally has irregularly spaced pitting on surface of drilled cores.
- Shale A fissile very fine-grained rock. Fractures along bedding planes.
- Limestone Rock made up predominantly of calcite (CaCO<sub>3</sub>). Effervesces strongly upon the application of dilute hydrochloric acid.
- Coal Rock consisting mainly of organic remains.
- Others Numerous other sedimentary rock types are present in lesser amounts in the stratigraphic record. The local abundance of any of these rock types is dependent upon the depositional history of the area. Conglomerate, halite, gypsum, dolomite, anhydrite, lignite, etc. are some of the rock types found in lesser amounts.

In classifying a sedimentary rock the following hierarchy shall be noted:

- Rock type
- Color
- Bedding thickness
- Hardness
- Fracturing
- Weathering
- Other characteristics

# 5.3.1 Rock Type

As described above, there are numerous types of sedimentary rocks. In most cases, a rock will be a combination of several grain types, therefore, a modifier such as a sandy siltstone, or a silty sandstone can be used. The modifier indicates that a significant portion of the rock type is composed of the modifier. Other modifiers can include carbonaceous, calcareous, siliceous, etc.

Grain size is the basis for the classification of clastic sedimentary rocks. Figure 4 is the Udden-Wentworth classification that will be assigned to sedimentary rocks. The individual boundaries are slightly different than the USCS subdivision for soil classification. For field determination of grain sizes, a scale can be used for the coarse grained rocks. For example, the division between siltstone and claystone may not be measurable in the field. The boundary shall be determined by use of a hand lens. If the grains cannot be seen with the naked eye but are distinguishable with a hand lens, the rock is a siltstone. If the grains are not distinguishable with a hand lens, the rock is a claystone.

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FIGURE 4

GRAIN SIZE CLASSIFICATION FOR ROCKS

Particle Name	Grain Size Diameter		
Cobbles	> 64 mm		
Pebbles	4 - 64 mm		
Granules	2 - 4 mm		
Very Coarse Sand	1 - 2 mm		
Coarse Sand	0.5 - 1 mm		
Medium Sand	0.25 - 0.5 mm		
Fine Sand	0.125 - 0.25 mm		
Very Fine Sand	0.0625 - 0.125 mm		
Silt	0.0039 - 0.0625 mm		

After Wentworth, 1922

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#### 5.3.2 Color

The color of a rock can be determined in a similar manner as for soil samples. Rock core samples shall be classified while wet, when possible, and air cored samples shall be scraped clean of cuttings prior to color classifications.

Rock color charts shall not be used unless specified by the Project Manager.

# 5.3.3 Bedding Thickness

The bedding thickness designations applied to soil classification (see Figure 3) will also be used for rock classification.

#### 5.3.4 Hardness

The hardness of a rock is a function of the compaction, cementation, and mineralogical composition of the rock. A relative scale for sedimentary rock hardness is as follows:

- Soft Weathered, considerable erosion of core, easily gouged by screwdriver, scratched by fingernail.
   Soft rock crushes or deforms under pressure of a pressed hammer. This term is always used for the hardness of the saprolite (decomposed rock which occupies the zone between the lowest soil horizon and firm bedrock).
- Medium soft Slight erosion of core, slightly gouged by screwdriver, or breaks with crumbly edges from single hammer blow.
- Medium hard No core erosion, easily scratched by screwdriver, or breaks with sharp edges from single hammer blow.
- Hard Requires several hammer blows to break and has sharp conchoidal breaks. Cannot be scratched with screwdriver.

Note the difference in usage here of the works "scratch" and "gouge." A scratch shall be considered a slight depression in the rock (do not mistake the scraping off of rock flour from drilling with a scratch in the rock itself), while a gouge is much deeper.

# 5.3.5 Fracturing

The degree of fracturing or brokenness of a rock is described by measuring the fractures or joint spacing. After eliminating drilling breaks, the average spacing is calculated and the fracturing is described by the following terms:

- Very broken (V. BR.) Less than 2-inch spacing between fractures
- Broken (BR.) 2-inch to 1-foot spacing between fractures
- Blocky (BL.) 1- to 3-foot spacing between fractures
- Massive (M.) 3 to 10-foot spacing between fractures

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The structural integrity of the rock can be approximated by calculating the Rock Quality Designation (RQD) of cores recovered. The RQD is determined by adding the total lengths of all pieces exceeding 4 inches and dividing by the total length of the coring run, to obtain a percentage.

Method of Calculating RQD (After Deere, 1964)

 $RQD \% = r/l \times 100$ 

- r = Total length of all pieces of the lithologic unit being measured, which are greater than 4 inches length, and have resulted from natural breaks. Natural breaks include slickensides, joints, compaction slicks, bedding plane partings (not caused by drilling), friable zones, etc.
- I = Total length of the coring run.

# 5.3.6 Weathering

The degree of weathering is a significant parameter that is important in determining weathering profiles and is also useful in engineering designs. The following terms can be applied to distinguish the degree of weathering:

- Fresh Rock shows little or no weathering effect. Fractures or joints have little or no staining and rock has a bright appearance.
- Slight Rock has some staining which may penetrate several centimeters into the rock. Clay filling of joints may occur. Feldspar grains may show some alteration.
- Moderate Most of the rock, with exception of quartz grains, is stained. Rock is weakened due to weathering and can be easily broken with hammer.
- Severe All rock including quartz grains is stained. Some of the rock is weathered to the extent of becoming a soil. Rock is very weak.

#### 5.3.7 Other Characteristics

The following items shall be included in the rock description:

- Description of contact between two rock units. These can be sharp or gradational.
- Stratification (parallel, cross stratified).
- Description of any filled cavities or vugs.
- Cementation (calcareous, siliceous, hematitic).
- Description of any joints or open fractures.
- Observation of the presence of fossils.
- Notation of joints with depth, approximate angle to horizontal, any mineral filling or coating, and degree of weathering.

All information shown on the boring logs shall be neat to the point where it can be reproduced on a copy machine for report presentation. The data shall be kept current to provide control of the drilling program and to indicate various areas requiring special consideration and sampling.

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# 5.3.8 Additional Terms Used in the Description of Rock

The following terms are used to further identify rocks:

- Seam Thin (12 inches or less), probably continuous layer.
- Some Indicates significant (15 to 40 percent) amounts of the accessory material. For example, rock composed of seams of sandstone (70 percent) and shale (30 percent) would be "sandstone -- some shale seams."
- Few Indicates insignificant (0 to 15 percent) amounts of the accessory material. For example, rock composed of seam of sandstone (90 percent) and shale (10 percent) would be "sandstone -- few shale seams."
- Interbedded Used to indicate thin or very thin alternating seams of material occurring in approximately equal amounts. For example, rock composed of thin alternating seams of sandstone (50 percent) and shale (50 percent) would be "interbedded sandstone and shale."
- Interlayered Used to indicate thick alternating seams of material occurring in approximately equal amounts.

The preceding sections describe the classification of sedimentary rocks. The following are some basic names that are applied to igneous rocks:

- Basalt A fine-grained extrusive rock composed primarily of calcic plagioclase and pyroxene.
- Rhyolite A fine-grained volcanic rock containing abundant quartz and orthoclase. The fine-grained equivalent of a granite.
- Granite A coarse-grained plutonic rock consisting essentially of alkali feldspar and quartz.
- Diorite A coarse-grained plutonic rock consisting essentially of sodic plagioclase and hornblende.
- Gabbro A coarse-grained plutonic rock consisting of calcic plagioclase and clinopyroxene. Loosely used for any coarse-grained dark igneous rock.

The following are some basic names that are applied to metamorphic rocks:

- Slate A very fine-grained foliated rock possessing a well developed slaty cleavage. Contains predominantly chlorite, mica, quartz, and sericite.
- Phyllite A fine-grained foliated rock that splits into thin flaky sheets with a silky sheen on cleavage surface.
- Schist A medium to coarse-grained foliated rock with subparallel arrangement of the micaceous minerals which dominate its composition.
- Gneiss A coarse-grained foliated rock with bands rich in granular and platy minerals.
- Quartzite A fine- to coarse-grained nonfoliated rock breaking across grains, consisting essentially of quartz sand with silica cement.

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# 5.4 Abbreviations

Abbreviations may be used in the description of a rock or soil. However, they shall be kept at a minimum. Following are some of the abbreviations that may be used:

С	-	Coarse	Lt	-	Light	YI	-	Yellow
Med	-	Medium	BR	-	Broken	Or	-	Orange
F	-	Fine	BL	-	Blocky	SS	-	Sandstone
V	-	Very	М	-	Massive	Sh	-	Shale
SI	-	Slight	Br	-	Brown	LS	-	Limestone
Осс	-	Occasional	ВІ	-	Black	Fgr	-	Fine-grained
Tr	-	Trace						

# 5.5 Boring Logs and Documentation

This section describes in more detail the procedures to be used in completing boring logs in the field. Information obtained from the preceding sections shall be used to complete the logs. A sample boring log has been provided as Figure 5.

The field geologist/engineer shall use this example as a guide in completing each boring log. Each boring log shall be fully described by the geologist/engineer as the boring is being drilled. Every sheet contains space for 25 feet of log. Information regarding classification details is provided either on the back of the boring log or on a separate sheet, for field use.

# 5.5.1 Soil Classification

- Identify site name, boring number, job number, etc. Elevations and water level data to be entered when surveyed data is available.
- Enter sample number (from SPT) under appropriate column. Enter depth sample was taken from (1 block = 1 foot). Fractional footages, i.e., change of lithology at 13.7 feet, shall be lined off at the proportional location between the 13- and 14-foot marks. Enter blow counts (Standard Penetration Resistance) diagonally (as shown). Standard penetration resistance is covered in Section 5.2.3.
- Determine sample recovery/sample length as shown. Measure the total length of sample recovered from the split-spoon sampler, including material in the drive shoe. Do not include cuttings or wash material that may be in the upper portion of the sample tube.
- Indicate any change in lithology by drawing a line at the appropriate depth. For example, if clayey silt
  was encountered from 0 to 5.5 feet and shale from 5.5 to 6.0 feet, a line shall be drawn at this
  increment. This information is helpful in the construction of cross-sections. As an alternative,
  symbols may be used to identify each change in lithology.
- The density of granular soils is obtained by adding the number of blows for the last two increments.
  Refer to Density of Granular Soils Chart on back of log sheet. For consistency of cohesive soils refer
  also to the back of log sheet Consistency of Cohesive Soils. Enter this information under the
  appropriate column. Refer to Section 5.2.3.

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FIGURE 5 COMPLETED BORING LOG (EXAMPLE)																	
BORING LOG Page 1 of 1																	
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- Enter color of the material in the appropriate column.
- Describe material using the USCS. Limit this column for sample description only. The predominant material is described last. If the primary soil is silt but has fines (clay) - use clayey silt. Limit soil descriptors to the following:

Trace: 0 - 10 percent
 Some: 11 - 30 percent
 And/Or: 31 - 50 percent

- Also indicate under Material Classification if the material is fill or natural soils. Indicate roots, organic material, etc.
- Enter USCS symbol use chart on back of boring log as a guide. If the soils fall into one of two basic groups, a borderline symbol may be used with the two symbols separated by a slash. For example ML/CL or SM/SP.
- The following information shall be entered under the "Remarks" column and shall include, but is not limited by, the following:
  - Moisture estimate moisture content using the following terms dry, moist, wet and saturated. These terms are determined by the individual. Whatever method is used to determine moisture, be consistent throughout the log.
  - Angularity describe angularity of coarse grained particles using the terms angular, subangular, subrounded, or rounded. Refer to ASTM D 2488 or Earth Manual for criteria for these terms.
  - Particle shape flat, elongated, or flat and elongated.
  - Maximum particle size or dimension.
  - Water level observations.
  - Reaction with HCI none, weak, or strong.
- Additional comments:
  - Indicate presence of mica, caving of hole, when water was encountered, difficulty in drilling, loss or gain of water.
  - Indicate odor and Photoionization Detector (PID) or Flame Ionization Detector (FID) reading if applicable.
  - Indicate any change in lithology by drawing a line through the lithology change column and indicate the depth. This will help when cross-sections are subsequently constructed.
  - At the bottom of the page indicate type of rig, drilling method, hammer size and drop, and any other useful information (i.e., borehole size, casing set, changes in drilling method).

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- Vertical lines shall be drawn (as shown in Figure 5) in columns 6 to 8 from the bottom of each sample to the top of the next sample to indicate consistency of material from sample to sample, if the material is consistent. Horizontal lines shall be drawn if there is a change in lithology, then vertical lines drawn to that point.
- Indicate screened interval of well, as needed, in the lithology column. Show top and bottom of screen. Other details of well construction are provided on the well construction forms.

#### 5.5.2 Rock Classification

- Indicate depth at which coring began by drawing a line at the appropriate depth. Indicate core run depths by drawing coring run lines (as shown) under the first and fourth columns on the log sheet. Indicate RQD, core run number, RQD percent, and core recovery under the appropriate columns.
- Indicate lithology change by drawing a line at the appropriate depth as explained in Section 5.5.1.
- Rock hardness is entered under designated column using terms as described on the back of the log or as explained earlier in this section.
- Enter color as determined while the core sample is wet; if the sample is cored by air, the core shall be scraped clean prior to describing color.
- Enter rock type based on sedimentary, igneous or metamorphic. For sedimentary rocks use terms as described in Section 5.3. Again, be consistent in classification. Use modifiers and additional terms as needed. For igneous and metamorphic rock types use terms as described in Sections 5.3.8.
- Enter brokenness of rock or degree of fracturing under the appropriate column using symbols VBR, BR, BL, or M as explained in Section 5.3.5 and as noted on the back of the Boring Log.
- The following information shall be entered under the remarks column. Items shall include but are not limited to the following:
  - Indicate depths of joints, fractures and breaks and also approximate to horizontal angle (such as high, low), i.e., 70° angle from horizontal, high angle.
  - Indicate calcareous zones, description of any cavities or vugs.
  - Indicate any loss or gain of drill water.
  - Indicate drop of drill tools or change in color of drill water.
- Remarks at the bottom of Boring Log shall include:
  - Type and size of core obtained.
  - Depth casing was set.
  - Type of rig used.
- As a final check the boring log shall include the following:
  - Vertical lines shall be drawn as explained for soil classification to indicate consistency of bedrock material.
  - If applicable, indicate screened interval in the lithology column. Show top and bottom of screen. Other details of well construction are provided on the well construction forms.

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# 5.5.3 Classification of Soil and Rock from Drill Cuttings

The previous sections describe procedures for classifying soil and rock samples when cores are obtained. However, some drilling methods (air/mud rotary) may require classification and borehole logging based on identifying drill cuttings removed from the borehole. Such cuttings provide only general information on subsurface lithology. Some procedures that shall be followed when logging cuttings are:

- Obtain cutting samples at approximately 5-foot intervals, sieve the cuttings (if mud rotary drilling) to
  obtain a cleaner sample, place the sample into a small sample bottle or "zip lock" bag for future
  reference, and label the jar or bag (i.e. hole number, depth, date, etc.). Cuttings shall be closely
  examined to determine general lithology.
- Note any change in color of drilling fluid or cuttings, to estimate changes in lithology.
- Note drop or chattering of drilling tools or a change in the rate of drilling, to determine fracture locations or lithologic changes.
- Observe loss or gain of drilling fluids or air (if air rotary methods are used), to identify potential fracture zones.
- Record this and any other useful information onto the boring log as provided in Figure 1.

This logging provides a general description of subsurface lithology and adequate information can be obtained through careful observation of the drilling process. It is recommended that split-barrel and rock core sampling methods be used at selected boring locations during the field investigation to provide detailed information to supplement the less detailed data generated through borings drilled using air/mud rotary methods.

# 5.6 Review

Upon completion of the borings logs, copies shall be made and reviewed. Items to be reviewed include:

- Checking for consistency of all logs.
- Checking for conformance to the guideline.
- Checking to see that all information is entered in their respective columns and spaces.

#### 6.0 REFERENCES

Unified Soil Classification System (USCS).

ASTM D2488, 1985.

Earth Manual, U.S. Department of the Interior, 1974.

#### 7.0 RECORDS

Originals of the boring logs shall be retained in the project files.



TETRA TECH NUS, INC.

# STANDARD OPERATING PROCEDURES

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Applicability

Tetra Tech NUS, Inc.

Prepared

Health & Safety

Subject

UTILITY LOCATING AND EXCAVATION CLEARANCE

Approved D. Senovich

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#### 1.0 PURPOSE

Utilities such as electric service lines, natural or propane gas lines, water and sewage lines, telecommunications, and steam lines are very often in the immediate vicinity of work locations. Contact with underground or overhead utilities can have serious consequences including employee injury/fatality, property and equipment damage, substantial financial impacts, and loss of utility service to users.

The purpose of this procedure is to provide minimum requirements and technical guidelines regarding the appropriate procedures to be followed when performing subsurface and overhead utility locating services. It is the policy of Tetra Tech NUS, Inc. (TtNUS) to provide a safe and healthful work environment for the protection of our employees. The purpose of this Standard Operating Procedure (SOP) is to aid in achieving the objectives of this policy, to present the acceptable procedures pertaining to utility locating and excavation clearance activities, and to present requirements and restrictions relevant to these types of activities. This SOP must be reviewed by any employee potentially involved with underground or overhead utility locating and avoidance activities.

## 2.0 SCOPE

This procedure applies to all TtNUS field activities where there may be potential contact with underground or overhead utilities. This procedure provides a description of the principles of operation, instrumentation, applicability, and implementability of typical methods used to determine the presence and avoidance of contact with utility services. This procedure is intended to assist with work planning and scheduling, resource planning, field implementation, and subcontractor procurement. Utility locating and excavation clearance requires site-specific information prior to the initiation of any such activities on a specific project. This SOP is not intended to provide a detailed description of methodology and instrument operation. Specialized expertise during both planning and execution of several of the methods presented may also be required.

#### 3.0 GLOSSARY

<u>Electromagnetic Induction (EMI) Survey</u> - A geophysical exploration method whereby electromagnetic fields are induced in the ground and the resultant secondary electromagnetic fields are detected as a measure of ground conductivity.

Magnetometer - A device used for precise and sensitive measurements of magnetic fields.

 $\underline{\text{Magnetic Survey}} - A$  geophysical survey method that depends on detection of magnetic anomalies caused by the presence of buried ferromagnetic objects.

<u>Metal Detection</u> – A geophysical survey method that is based on electromagnetic coupling caused by underground conductive objects.

<u>Vertical Gradiometer</u> – A magnetometer equipped with two sensors that are vertically separated by a fixed distance. It is best suited to map near surface features and is less susceptible to deep geologic features.

<u>Ground Penetrating Radar</u> – Ground Penetrating Radar (GPR) involves specialized radar equipment whereby a signal is sent into the ground via a transmitter. Some portion of the signal will be reflected from the subsurface material, which is then recorded with a receiver and electronically converted into a graphic picture.

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#### 4.0 RESPONSIBILITIES

<u>Project Manager (PM)/Task Order Manager (TOM)</u> - Responsible for ensuring that all field activities are conducted in accordance with this procedure.

<u>Site Manager (SM)/Field Operations Leader (FOL)</u> - Responsible for the onsite verification that all field activities are performed in compliance with approved SOPs or as otherwise directed by the approved project plan(s).

<u>Site Health & Safety Officer (SHSO)</u> – Responsible to provide technical assistance and verify full compliance with this SOP. The SHSO is also responsible for reporting any deficiencies to the Corporate Health and Safety Manager (HSM) and to the PM/TOM.

<u>Health & Safety Manager (HSM)</u> – Responsible for preparing, implementing, and modifying corporate health and safety policy and this SOP.

<u>Site Personnel</u> – Responsible for performing their work activities in accordance with this SOP and the TtNUS Health and Safety Policy.

#### 5.0 PROCEDURES

This procedure addresses the requirements and technical procedures that must be performed to minimize the potential for contact with underground and overhead utility services. These procedures are addressed individually from a buried and overhead standpoint.

## 5.1 Buried Utilities

Buried utilities present a heightened concern because their location is not typically obvious by visual observation, and it is common that their presence and/or location is unknown or incorrectly known on client properties. This procedure must be followed prior to beginning any subsurface probing or excavation that might potentially be in the vicinity of underground utility services. In addition, the Utility Clearance Form (Attachment 3) must be completed for every location or cluster of locations where intrusive activities will occur.

Where the positive identification and de-energizing of underground utilities cannot be obtained and confirmed using the following steps, the PM/TOM is responsible for arranging for the procurement of a qualified, experienced, utility locating subcontractor who will accomplish the utility location and demarcation duties specified herein.

- 1. A comprehensive review must be made of any available property maps, blue lines, or as-builts prior to site activities. Interviews with local personnel familiar with the area should be performed to provide additional information concerning the location of potential underground utilities. Information regarding utility locations shall be added to project maps upon completion of this exercise.
- 2., A visual site inspection must be performed to compare the site plan information to actual field conditions. Any findings must be documented and the site plan/maps revised. The area(s) of proposed excavation or other subsurface activities must be marked at the site in white paint or pin flags to identify those locations of the proposed intrusive activities. The site inspection should focus on locating surface indications of potential underground utilities. Items of interest include the presence of nearby area lights, telephone service, drainage grates, fire hydrants, electrical service vaults/panels, asphalt/concrete scares and patches, and topographical depressions. Note the location of any emergency shut off switches. Any additional information regarding utility

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locations shall be added to project maps upon completion of this exercise and returned to the PM/TOM.

- 3. If the planned work is to be conducted on private property (e.g., military installations, manufacturing facilities, etc.) the FOL must identify and contact appropriate facility personnel (e.g., public works or facility engineering) before any intrusive work begins to inquire about (and comply with) property owner requirements. It is important to note that private property owners may require several days to several weeks advance notice prior to locating utilities.
- 4. If the work location is on public property, the state agency that performs utility clearances must be notified (see Attachment 1). State "one-call" services must be notified prior to commencing fieldwork per their requirements. Most one-call services require, by law, 48- to 72-hour advance notice prior to beginning any excavation. Such services typically assign a "ticket" number to the particular site. This ticket number must be recorded for future reference and is valid for a specific period of time, but may be extended by contacting the service again. The utility service will notify utility representatives who then mark their respective lines within the specified time frame. It should be noted that most military installations own their own utilities but may lease service and maintenance from area providers. Given this situation, "one call" systems may still be required to provide location services on military installations.
- 5. Utilities must be identified and their locations plainly marked using pin flags, spray paint, or other accepted means. The location of all utilities must be noted on a field sketch for future inclusion on project maps. Utility locations are to be identified using the following industry-standard color code scheme, unless the property owner or utility locator service uses a different color code:

white excavation/subsurface investigation location

red electrical yellow gas, oil, steam

orange telephone, communications

blue water, irrigation, slurry

green sewer, drain

- 6. Where utility locations are not confirmed with a high degree of confidence through drawings, schematics, location services, etc., the work area must be thoroughly investigated prior to beginning the excavation. In these situations, utilities must be identified using safe and effective methods such as passive and intrusive surveys, or the use of non-conductive hand tools. Also, in situations where such hand tools are used, they should always be used in conjunction with suitable detection equipment, such as the items described in Section 6.0 of this SOP. Each method has advantages and disadvantages including complexity, applicability, and price. It also should be noted that in some states, initial excavation is required by hand to a specified depth.
- 7. At each location where trenching or excavating will occur using a backhoe or other heavy equipment, and where utility identifications and locations cannot be confirmed prior to groundbreaking, the soil must be probed using a device such as a tile probe which is made of non-conductive material such as fiberglass. If these efforts are not successful in clearing the excavation area of suspect utilities, hand shoveling must be performed for the perimeter of the intended excavation.
- 8. All utilities uncovered or undermined during excavation must be structurally supported to prevent potential damage. Unless necessary as an emergency corrective measure, TtNUS shall not make any repairs or modifications to existing utility lines without prior permission of the utility owner, property owner, and Corporate HSM. All repairs require that the line be locked-out/tagged-out prior to work.

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# 5.2 Overhead Power Lines

If it is necessary to work within the minimum clearance distance of an overhead power line, the overhead line must be de-energized and grounded, or re-routed by the utility company or a registered electrician. If protective measures such as guarding, isolating, or insulating are provided, these precautions must be adequate to prevent employees from contacting such lines directly with any part of their body or indirectly though conductive materials, tools, or equipment.

The following table provides the required minimum clearances for working in proximity to overhead power lines.

Nominal Voltage	Minimum Clearance
0 -50 kV	10 feet, or one mast length; whichever is greater
50+ kV	10 feet plus 4 inches for every 10 kV over 50 kV or 1.5
	mast lengths; whichever is greater

## 6.0 UNDERGROUND LOCATING TECHNIQUES

A variety of supplemental utility locating approaches are available and can be applied when additional assurance is needed. The selection of the appropriate method(s) to employ is site-specific and should be tailored to the anticipated conditions, site and project constraints, and personnel capabilities.

## 6.1 Geophysical Methods

Geophysical methods include electromagnetic induction, magnetics, and ground penetrating radar. Additional details concerning the design and implementation of electromagnetic induction, magnetics, and ground penetrating radar surveys can be found in one or more of the TtNUS SOPs included in the References (Section 8.0).

## **Electromagnetic Induction**

Electromagnetic Induction (EMI) line locators operate either by locating a background signal or by locating a signal introduced into the utility line using a transmitter. A utility line acts like a radio antenna, producing electrons, which can be picked up with a radiofrequency receiver. Electrical current carrying conductors have a 60HZ signal associated with them. This signal occurs in all power lines regardless of voltage. Utilities in close proximity to power lines or used as grounds may also have a 60HZ signal, which can be picked up with an EM receiver. A typical example of this type of geophysical equipment is an EM-61.

EMI locators specifically designed for utility locating use a special signal that is either indirectly induced onto a utility line by placing the transmitter above the line or directly induced using an induction clamp. The clamp induces a signal on the specific utility and is the preferred method of tracing since there is little chance of the resulting signals being interfered with. A good example of this type of equipment is the Schonstedt® MAC-51B locator. The MAC-51B performs inductively traced surveys, simple magnetic locating, and traced nonmetallic surveys.

When access can be gained inside a conduit to be traced, a flexible insulated trace wire can be used. This is very useful for non-metallic conduits but is limited by the availability of gaining access inside the pipe.

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## Magnetics

Magnetic locators operate by detecting the relative amounts of buried ferrous metal. They are incapable of locating or identifying nonferrous utility lines but can be very useful for locating underground storage tanks (UST's), steel utility lines, and buried electrical lines. A typical example of this type of equipment is the Schonstedt® GA-52Cx locator. The GA-52Cx is capable of locating 4-inch steel pipe up to 8 feet deep.

Non-ferrous lines are often located by using a typical plumbing tool (snake) fed through the line. A signal is then introduced to the snake that is then traced.

# **Ground Penetrating Radar**

Ground Penetrating Radar (GPR) involves specialized radar equipment whereby a signal is sent into the ground via a transmitter. Some portion of the signal will be reflected from the subsurface material, which is then recorded with a receiver and electronically converted into a graphic picture. In general, an object which is harder than the surrounding soil will reflect a stronger signal. Utilities, tunnels, UST's, and footings will reflect a stronger signal than the surrounding soil. Although this surface detection method may determine the location of a utility, this method does not specifically identify utilities (i.e., water vs. gas, electrical vs. telephone); hence, verification may be necessary using other methods. This method is somewhat limited when used in areas with clay soil types or with a high water table.

## 6.2 Passive Detection Surveys

#### **Acoustic Surveys**

Acoustic location methods are generally most applicable to waterlines or gas lines. A highly sensitive Acoustic Receiver listens for background sounds of water flowing (at joints, leaks, etc.) or to sounds introduced into the water main using a transducer. Acoustics may also be applicable to determine the location of plastic gas lines.

## Thermal Imaging

Thermal (i.e., infrared) imaging is a passive method for detecting the heat emitted by an object. Electronics in the infrared camera convert subtle heat differentials into a visual image on the viewfinder or a monitor. The operator does not look for an exact temperature; rather they look for heat anomalies (either elevated or suppressed temperatures) characteristic of a potential utility line.

The thermal fingerprint of underground utilities results from differences in temperature between the atmosphere and the fluid present in a pipe or the heat generated by electrical resistance. In addition, infrared scanners may be capable of detecting differences in the compaction, temperature and moisture content of underground utility trenches. High-performance thermal imagery can detect temperature differences to hundredths of a degree.

# 6.3 <u>Intrusive Detection Surveys</u>

#### **Vacuum Excavation**

Vacuum excavation is used to physically expose utility services. The process involves removing the surface material over approximately a 1' x 1' area at the site location. The air-vacuum process proceeds with the simultaneous action of compressed air-jets to loosen soil and vacuum extraction of the resulting

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debris. This process ensures the integrity of the utility line during the excavation process, as no hammers, blades, or heavy mechanical equipment comes into contact with the utility line, eliminating the risk of damage to utilities. The process continues until the utility is uncovered. Vacuum excavation can be used at the proposed site location to excavate below the "utility window" which is usually 8 feet.

#### Hand Excavation

When the identification and location of underground utilities cannot be positively confirmed through document reviews and/or other methods, borings and excavations may be cleared via the use of nonconductive hand tools. This should always be done in conjunction with the use of detection equipment. This would be required for all locations where there is a potential to impact buried utilities. The minimum hand-excavation depth that must be reached is to be determined considering the geographical location of the work site. This approach recognizes that the placement of buried utilities is influenced by frost line depths that vary by geographical region. Attachment 2 presents frost line depths for the regions of the contiguous United States. At a minimum, hand excavation depths must be at least to the frost line depth (see Attachment 2) plus two (2) feet, but never less than 4 feet below ground surface (bgs). For hand excavation, the hole created must be reamed large enough to be at least the diameter of the drill rig auger or bit prior to drilling. For soil gas surveys, the survey probe shall be placed as close as possible to the cleared hand excavation. It is important to note that a post-hole digger must not be used in this type of hand excavation activity.

# **Tile Probe Surveys**

For some soil types, site conditions, and excavation requirements, non-conductive tile probes may be used. A tile probe is a "T"-handled rod of varying lengths that can be pushed into the soil to determine if any obstructions exist at that location. Tile probes constructed of fiberglass or other nonconductive material are readily-available from numerous vendors. Tile probes must be performed to the same depth requirements as previously specified. As with other types of hand excavating activities, the use of a nonconductive tile probe, should always be in conjunction with suitable utility locating detection equipment.

#### 7.0 INTRUSIVE ACTIVITIES SUMMARY

The following list summarizes the activities that must be performed prior to beginning subsurface activities:

- 1. Map and mark all subsurface locations and excavation boundaries using white paint or markers specified by the client or property owner.
- 2. Notify the property owner and/or client that the locations are marked. At this point, drawings of locations or excavation boundaries shall be provided to the property owner and/or client so they may initiate (if applicable) utility clearance.
  - Note: Drawings with confirmed locations should be provided to the property owner and/or client as soon as possible to reduce potential time delays.
- 3. Notify "One Call" service. If possible, arrange for an appointment to show the One Call representative the surface locations or excavation boundaries in person. This will provide a better location designation to the utilities they represent. You should have additional drawings should you need to provide plot plans to the One Call service.
- 4. Implement supplemental utility detection techniques as necessary and appropriate to conform utility locations or the absence thereof.

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5. Complete Attachment 3, Utility Clearance Form. This form should be completed for each excavation location. In situations where multiple subsurface locations exist within the close proximity of one another, one form may be used for multiple locations provided those locations are noted on the Utility Clearance Form. Upon completion, the Utility Clearance Form and revised/annotated utility location map becomes part of the project file.

## 8.0 REFERENCES

OSHA Letter of Interpretation, Mr. Joseph Caldwell, Attachment 4 OSHA 29 CFR 1926(b)(2) OSHA 29 CFR 1926(b)(3) TtNUS Utility Locating and Clearance Policy TtNUS SOP GH-3.1; Resistivity and Electromagnetic Induction TtNUS SOP GH-3.2; Magnetic and Metal Detection Surveys

TtNUS SOP GH-3.4; Ground-penetrating Radar Surveys

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# **ATTACHMENT 1** LISTING OF UNDERGROUND UTILITY CLEARANCE RESOURCES



American Public Works Association 2345 Grand Boulevard, Suite 500, Kansas City, MO 64108-2625 Phone (816) 472-6100 • Fax (816) 472-1610 Web www.apwa.net . E-mail apwa@apwa.net

#### **ONE-CALL SYSTEMS INTERNATIONAL CONDENSED DIRECTORY**

Alabama

Alabama One-Call 1-800-292-8525

Locate Call Center of Alaska, Inc. 1-800-478-3121

Arizona

Arizona Blue Stake 1-800-782-5348

Arkansas One Call System, Inc. 1-800-482-8998

California

Underground Service Alert North 1-800-227-2600 Underground Service Alert of Southern California 1-800-227-2600

Colorado

**Utility Notification Center of Colorado** 1-800-922-1987

Connecticut Call Before You Dig 1-800-922-4455

Miss Utility of Delmarva 1-800-282-8555

Sunshine State One-Call of Florida, Inc. 1-800-432-4770

Underground Protection Center, Inc. 1-800-282-7411

Hawali

Underground Service Alert North 1-800-227-2600

Idaho

Dig Line Inc. 1-800-342-1585 Kootenal County One-Call 1-800-428-4950 Shoshone - Benewah One-Call 1-800-398-3285

Illinois

JULIE, Inc. 1-800-892-0123 Digger (Chicago Utility Alert Network) 312-744-7000

Indiana

Indiana Underground Plant Protection Service 1-800-382-5544

Iowa One-Call 1-800-292-8989

Kansas Kansas One-Call System, Inc.

1-800-344-7233

Kentucky

Kentucky Underground Protection Inc. 1-800-752-6007

Louisiana One Call System, Inc. 1-800-272-3020

Maine

Dig Safe System, Inc. 1-888-344-7233

Marviand

Miss Utility 1-800-257-7777 Miss Utility of Delmarva 1-800-282-8555

Massachusetts

Dig Safe System, Inc. 1-888-344-7233

Michigan

Miss Dig System, Inc. 1-800-482-7171

Minnesota

Gopher State One Call 1-800-252-1168

Mississippi

Mississippi One-Call System, Inc. 1-800-227-6477

Missouri

Missouri One-Call System, Inc. 1-800-344-7483

Montana

Utilities Underground Protection Center 1-800-424-5555 Montana One Call Center 1-800-551-8344

Nebraska

Diggers Hotline of Nebraska

1-800-331-5666

Underground Service Afert North 1-800-227-2600

New Hampshire Dig Safe System, Inc. 1-888-344-7233 New Jersey

New Jersey One Call 1-800-272-1000

**New Mexico** 

New Mexico One Call System, Inc. 1-800-321-2537 Las Cruces- Dona Ana Blue Stakes 1-888-526-0400

**New York** 

Dig Safely New York 1-800-962-7962 New York City- Long Island One Call Center 1-800-272-4480

North Carolina

The North Carolina One-Call Center, Inc. 1-800-632-4949

North Dakota North Dakota One-Call 1-800-795-0555

Ohio Utilities Protection Service 1-800-362-2764 Oil & Gas Producers Underground Protect'n Svc 1-800-925-0988

Oklahoma

Call Okie 1-800-522-6543

Oregon Utility Notification Center/One Call Concepts 1-800-332-2344

Pennsylvania

Pennsylvania One Call System, Inc. 1-800-242-1776

Rhode Island

Dig Safe System, Inc. 1-888-344-7233

South Carolina Palmetto Utility Protection Service Inc. 1-888-721-7877

South Dakota South Dakota One Cali

1-800-781-7474

Tennessee Tennessee One-Call System, Inc. 1-800-351-1111

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# **ATTACHMENT 1 (Continued)**

Texas

Texas One Call System 1-800-245-4545 Texas Excavation Safety System, Inc. 1-800-344-8377 Lone Star Notification Center 1-800-669-8344

Utah

Blue Stakes of Utah 1-800-662-4111

Dig Safe System, Inc. 1-888-344-7233

Virginia

Miss Utility of Virginia 1-800-552-7001 Miss Utility (Northern Virginia) 1-800-257-7777

Washington

**Utilities Underground Location Center** 1-800-424-5555 Northwest Utility Notification Center 1-800-553-4344 Inland Empire Utility Coordinating Council 509-456-8000

West Virginia Miss Utility of West Virginia, Inc. 1-800-245-4848

Wisconsin

Diggers Hotline, Inc. 1-800-242-8511

Wyoming One-Call System, Inc. 1-800-348-1030 Call Before You Dig of Wyoming 1-800-849-2476 District of Columbia

Miss Utility 1-800-257-7777

Alberta

Alberta One-Call Corporation 1-800-242-3447

**British Columbia** BC One Call 1-800-474-6886

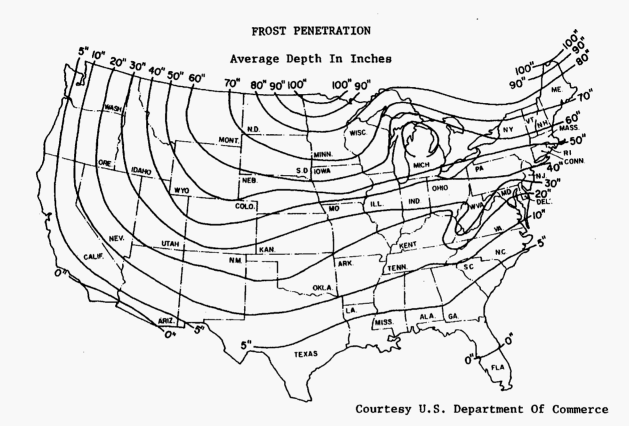
Ontario Ontario One-Call System 1-800-400-2255

Quebec Info-Excavation 1-800-663-9228

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# **ATTACHMENT 2**

# FROST LINE PENETRATION DEPTHS BY GEOGRAPHIC LOCATION



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# ATTACHMENT 3 UTILITY CLEARANCE FORM

t:	Project Name:	
ct No	: Completed By:	
vation	Method/Overhead Equipment:	
Ur	derground Utilities	<u>Circle One</u>
a)	Review of existing maps?	yes no N/A
b)	Interview local personnel?	yes no N/A
c)	Site visit and inspection?	yes no N/A
d)	Excavation areas marked in the field?	yes no N/A
e)	Utilities located in the field?	yes no N/A
f)	Located utilities marked/added to site maps?	yes no N/A
g)	Client contact notified	yes no N/A
	Name Telephone: Date:	
g)	State One-Call agency called?	yes no N/A
	Caller: Date:	
h)	Geophysical survey performed?	yes no N/A
	Survey performed by: Date:	
i)	Hand excavation performed (with concurrent use of utility	
'/	detection device)?	yes no NA
	Completed by:feet Date:	
j)	Trench/excavation probed?	— yes no N/A
J <i>)</i>	Probing completed by:	
	Depth/frequency: Date:	
O۱	erhead Utilities	Present Abser
a)	Determination of nominal voltage	yes no N/A
b) c)	Marked on site maps Necessary to lockout/insulate/re-route	yes no N/A yes no N/A
d)	Document procedures used to lockout/insulate/re-route	yes no N/A
e)	Minimum acceptable clearance (SOP Section 5.2):	
No	tes:	
_		
_		
Ap	proval:	
Sit	e Manager/Field Operations Leader Date	
		c: PM/Project Fi Program Fi

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# ATTACHMENT 4 OSHA LETTER OF INTERPRETATION

Mr. Joseph Caldwell Consultant Governmental Liaison Pipeline Safety Regulations 211 Wilson Boulevard Suite 700 Arlington, Virginia 22201

Re: Use of hydro-vacuum or non-conductive hand tools to locate underground utilities.

#### Dear Mr. Caldwell:

In a letter dated July 7, 2003, we responded to your inquiry of September 18, 2002, regarding the use of hydro-vacuum equipment to locate underground utilities by excavation. After our letter to you was posted on the OSHA website, we received numerous inquiries that make it apparent that aspects of our July 7 letter are being misunderstood. In addition, a number of industry stakeholders, including the National Utility Contractors Association (NUCA), have provided new information regarding equipment that is available for this work.

To clarify these issues, we are withdrawing our July 7 letter and issuing this replacement response to your inquiry.

Question: Section 1926.651 contains several requirements that relate to the safety of employees engaged in excavation work. Specifically, paragraphs (b)(2) and (b)(3) relate in part to the safety of the means used to locate underground utility installations that, if damaged during an uncovering operation, could pose serious hazards to employees.

Under these provisions, what constitutes an acceptable method of uncovering underground utility lines, and further, would the use of hydro-vacuum excavation be acceptable under the standard?

#### **Answer**

# Background

Two sections of 29 CFR 1926 Subpart P (Excavations), 1926.651(Specific excavation requirements), govern methods for uncovering underground utility installations. Specifically, paragraph (b)(2) states:

When utility companies or owners cannot respond to a request to locate underground utility installations within 24 hours \* \* \* or cannot establish the exact location of these installations, the employer may proceed, provided the employer does so with caution, and provided detection equipment or other acceptable means to locate utility installations are used. (emphasis added).

Paragraph (b)(3) provides:

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# **ATTACHMENT 4 (Continued)**

When excavation operations approach the estimated location of underground installations, the exact location of the installations shall be determined by <u>safe and acceptable means</u>. (emphasis added).

Therefore, "acceptable means" must be used where the location of the underground utilities have not been identified by the utility companies and detection equipment is not used.

Subpart P does not contain a definition of either "other acceptable means" or "safe and acceptable means." The preambles to both the proposed rule and the final rule discussed the rationale behind the wording at issue. For example, the preamble to the proposed rule, 52 Fed. Reg. 12301 (April 15, 1987), noted that a 1972 version of this standard contained language that specified "careful probing or hand digging" as the means to uncover utilities. The preamble then noted that an amendment to the 1972 standard later deleted that language "to allow other, equally effective means of locating such installations." The preamble continued that in the 1987 proposed rule, OSHA again proposed using language in section (b)(3) that would provide another example of an acceptable method of uncovering utilities that could be used where the utilities have not been marked and detection equipment is not being used—"probing with hand-held tools." This method was rejected in the final version of 29 CFR 1926. As OSHA explained in the preamble to the final rule, 54 Fed. Reg. 45916 (October 31, 1989):

OSHA received two comments \*\*\* and input from ACCSH [OSHA's Advisory Committee on Construction Safety and Health] \*\*\* on this provision. All commenters recommended dropping 'such as probing with hand-held tools' from the proposed provision, because this could create a hazard to employees by damaging the installation or its insulation.

In other words, the commenters objected to the use of hand tools being used unless detection equipment was used in conjunction with them. OSHA then concluded its discussion relative to this provision by agreeing with the commentators and ultimately not including any examples of "acceptable means" in the final provision.

# Non-conductive hand tools are permitted

This raises the question of whether the standard permits the use of hand tools alone -- without also using detection equipment. NUCA and other industry stakeholders have recently informed us that non-conductive hand tools that are appropriate to be used to locate underground utilities are now commonly available.

Such tools, such as a "shooter" (which has a non-conductive handle and a snub nose) and non-conductive or insulated probes were not discussed in the rulemaking. Since they were not considered at that time, they were not part of the class of equipment that was thought to be unsafe for this purpose. Therefore, we conclude that the use of these types of hand tools, when used with appropriate caution, is an "acceptable means" for locating underground utilities.

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# **ATTACHMENT 4 (Continued)**

## Hydro-vacuum excavation

It is our understanding that some hydro-vacuum excavation equipment can be adjusted to use a minimum amount of water and suction pressure. When appropriately adjusted so that the equipment will not damage underground utilities (especially utilities that are particularly vulnerable to damage, such as electrical lines), use of such equipment would be considered a "acceptable means" of locating underground utilities. However, if the equipment cannot be sufficiently adjusted, then this method would not be acceptable under the standard.

# Other technologies

We are not suggesting that these are the only devices that would be "acceptable means" under the standard. Industry stakeholders have informed us that there are other types of special excavation equipment designed for safely locating utilities as well.

We apologize for any confusion our July 7 letter may have caused. If you have further concerns or questions, please feel free to contact us again by fax at: U.S. Department of Labor, OSHA, Directorate of Construction, Office of Construction Standards and Compliance Assistance, fax # 202-693-1689. You can also contact us by mail at the above office, Room N3468, 200 Constitution Avenue, N.W., Washington, D.C. 20210, although there will be a delay in our receiving correspondence by mail.

Sincerely,

Russell B. Swanson, Director Directorate of Construction

NOTE: OSHA requirements are set by statute, standards and regulations. Our interpretation letters explain these requirements and how they apply to particular circumstances, but they cannot create additional employer obligations. This letter constitutes OSHA=s interpretation of the requirements discussed. Note that our enforcement guidance may be affected by changes to OSHA rules. Also, from time to time we update our guidance in response to new information. To keep apprised of such developments, you can consult OSHA's website at http://www.osha.gov.



# STANDARD OPERATING PROCEDURES

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Applicability

Tetra Tech NUS, Inc.

Prepared

Earth Sciences Department

Subject

SOIL SAMPLING

Approved

Tom Johnston



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## 1.0 PURPOSE

This Standard Operating Procedure (SOP) describes the procedures to be used to collect surface, near-surface, and subsurface soil samples. Additionally, it describes the methods for sampling of test pits and trenches to determine subsurface soil and rock conditions and for recovery of small-volume or bulk samples from pits.

#### 2.0 SCOPE

This document applies to the collection of surface, near-surface, and subsurface soil samples exposed through hand digging, hand augering, drilling, or machine excavating at hazardous substance sites for laboratory testing, onsite visual examination, and onsite testing.

#### 3.0 GLOSSARY

<u>Composite Sample</u> - A composite sample is a combination of more than one grab sample from various locations and/or depths and times that is homogenized and treated as one sample. This type of sample is usually collected when determination of an average waste concentration for a specific area is required. Composite samples shall not be collected for volatile organics analysis.

Confined Space - As stipulated in 29 Code of Federal Regulations (CFR) 1910.146, a confined space means a space that: (1) is large enough and so configured that an employee can bodily enter and perform assigned work; (2) has limited or restricted means for entry or exit (e.g., tanks, vessels, silos, storage bins, hoppers, vaults, pits, and excavations); and (3) is not designed for continuous employee occupancy. TtNUS considers all confined space as permit-required confined spaces.

Grab Sample - One sample collected at one location and at one specific time.

Hand Auger - A sampling device used to extract soil from the ground.

Representativeness – A qualitative description of the degree to which an individual sample accurately reflects population characteristics or parameter variations at a sampling point. It is therefore an important characteristic not only of assessment and quantification of environmental threats posed by the site, but also for providing information for engineering design and construction. Proper sample location selection and proper sample collection methods are important to ensure that a truly representative sample has been collected.

<u>Sample for Non-Volatile Analyses</u> - Includes all chemical parameters other than volatile organics (e.g., semivolatiles, pesticides/PCBs, metals, etc.) and those engineering parameters that do not require undisturbed soil for their analysis.

<u>Split-Barrel Sampler</u> - A steel tube, split in half lengthwise, with the halves held together by threaded collars at either end of the tube. Also called a split-spoon sampler, this device can be driven into resistant materials using a drive weight mounted in the drilling string. A standard split-barrel sampler is typically available in two common lengths, providing either 20-inch or 26-inch longitudinal clearance for obtaining 18-inch or 24-inch-long samples, respectively. These split-barrel samplers commonly range in size from 2 to 3.5 inches OD. The larger sizes are commonly used when a larger volume of sample material is required (see Attachment B).

<u>Test Pit and Trench</u> - Open, shallow excavations, typically rectangular (if a test pit) or longitudinal (if a trench), excavated to determine shallow subsurface conditions for engineering, geological, and soil chemistry exploration and/or sampling purposes. These pits are excavated manually or by machine (e.g., backhoe, clamshell, trencher, excavator, or bulldozer).

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<u>Thin-Walled Tube Sampler</u> - A thin-walled metal tube (also called a Shelby tube) used to recover relatively undisturbed soil samples. These tubes are available in various sizes, ranging from 2 to 5 inches outside diameter (OD) and from 18 to 54 inches in length.

#### 4.0 RESPONSIBILITIES AND PERSONNEL QUALIFICATIONS

<u>Project Manager</u> - The Project Manager is responsible for determining the sampling objectives, selecting proposed sampling locations, and selecting field procedures used in the collection of soil samples. Additionally, in consultation with other project personnel (geologist, hydrogeologist, etc.), the Project Manager establishes the need for test pits or trenches and determines their approximate locations and dimensions.

<u>Site Safety Officer (SSO)</u> - The SSO (or a qualified designee) is responsible for providing the technical support necessary to implement the project Health and Safety Plan. This will include (but not be limited to) performing air quality monitoring during sampling, boring, and excavation activities and to ensure that workers and offsite (downwind) individuals are not exposed to hazardous levels of airborne contaminants. The SSO/designee may also be required to advise the FOL on other safety-related matters regarding boring, excavation, and sampling, such as mitigative measures to address potential hazards from unstable trench walls, puncturing of drums or other hazardous objects, etc.

<u>Field Operations Leader (FOL)</u> - This individual is primarily responsible for the execution of the planning document containing the Sampling and Analysis Plan (SAP). This is accomplished through management of a field sampling team for the proper acquisition of samples. He or she is responsible for the supervision of onsite analyses; ensuring proper instrument calibration, care, and maintenance; sample collection and handling; the completion and accuracy of all field documentation; and making sure that custody of all samples obtained is maintained according to proper procedures. When appropriate and as directed by the FOL, such responsibilities may be performed by other qualified personnel (e.g., field technicians) where credentials and time permit. The FOL is responsible for finalizing the locations for collection of surface, near-surface, and subsurface (hand and machine borings, test pits/trenches) soil samples. He/she is ultimately responsible for the sampling and backfilling of boreholes, test pits, and trenches and for adherence to Occupational Safety and Health Administration (OSHA) regulations during these operations through self acquisition or through the management of a field team of samplers.

<u>Project Geologist/Sampler</u> - The project geologist/sampler is responsible for the proper acquisition of samples in accordance with this SOP and/or other project-specific documents. In addition, this individual is responsible for the completion of all required paperwork (e.g., sample log sheets, field notebook, boring logs, test pit logs, container labels, custody seals, and chain-of-custody forms) associated with the collection of those samples.

<u>Competent Person</u> - A Competent Person, as defined in 29 CFR 1929.650 of Subpart P - Excavations, means one who is capable of identifying existing and predictable hazards in the surroundings, or working conditions that are unsanitary, hazardous, or dangerous to employees, and who has authorization to take prompt corrective measures to eliminate them.

General personnel qualifications for groundwater sample collection and onsite water quality testing include the following:

- Occupational Safety and Health Administration (OSHA) 40-hour and applicable refresher training.
- Capability of performing field work under the expected physical and environmental (i.e., weather)
  conditions.

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• Familiarity with appropriate procedures for sample documentation, handling, packaging, and shipping.

#### 5.0 HEALTH AND SAFETY

Health and safety precautions are identified for individual sample collection procedures throughout this SOP. In addition to those precautions, the following general hazards may be incurred during sampling activities:

- Knee injuries from kneeling on hard or uneven surfaces
- Slips, trips, and falls
- Cuts and lacerations
- Traffic hazards associated with sampling in parking areas, along roadways and highways.

Methods of avoiding these hazards are provided below.

**Knee injuries** – If kneeling is required during soil sampling, this could result in knee injuries from stones/foreign objects and general damage due to stress on the joints. To minimize this hazard:

- Clear any foreign objects from the work area.
- Wear hard-sided knee pads.
- Stretch ligaments, tendons and muscles before, during and after. Take breaks as frequently as necessary.
- Report pre-existing conditions to the SSO if you feel this activity will aggravate an existing condition.

**Slips, Trips, and Falls** – These hazards exist while traversing varying terrains carrying equipment to sample locations. To minimize these hazards:

- Pre-survey sampling locations. Eliminate, barricade, or otherwise mark physical hazards leading to the locations.
- Carry small loads that do not restrict the field of vision.
- Travel the safest and clearest route (not necessarily the shortest).

**Cuts and Lacerations** - To prevent cuts and lacerations associated with soil sampling, the following provisions are required:

- Always cut away from yourself and others when cutting tubing or rope. This will prevent injury to yourself and others if the knife slips.
- Do not place items to be cut in your hand or on your knee.
- Change blades as necessary to maintain a sharp cutting edge. Many accidents result from struggling with dull cutting attachments.

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- Whenever practical, wear cut-resistant gloves (e.g., leather or heavy cotton work gloves) at least on the hand not using the knife.
- Keep cutting surfaces clean and smooth.
- Secure items to be cut do not hold them against the opposing hand, a leg, or other body part.
- When transporting glassware, keep it in a hard-sided container such as a cooler so that if there is a fall, you will be less likely to get cut by broken glass.
- DO NOT throw broken sample jars or glass ampoules into garbage bags. Place broken glass and glass ampoules in hard-sided containers such as a cardboard box or directly into a dumpster. DO NOT reach into garbage bags to retrieve any item accidentally thrown away. Empty the contents onto a flat surface to avoid punctures and lacerations from reaching where you cannot see.

**Vehicular and Foot Traffic Hazards** – When sampling along the roadway or near traffic patterns, follow the following precautions:

- Motorists may be distracted by onsite activities ASSUME THEY DO NOT SEE YOU OR MEMBERS
  OF YOUR FIELD CREW.
- DO NOT place obstructions (such as vehicles) along the sides of the road that may cause site
  personnel to move into the flow of traffic to avoid your activities or equipment or that will create a
  blind spot.
- Provide a required free space of travel. Maintain at least 6 feet of space between you and moving traffic. Where this is not possible, use flaggers and/or signs to warn oncoming traffic of activities near or within the travel lanes.
- Face Traffic. Whenever feasible, if you must move within the 6 feet of the required free space or into traffic, attempt to face moving traffic at all times. Always leave yourself an escape route.
- Wear high-visibility vests to increase visual recognition by motorists.
- Do not rely on the vehicle operator's visibility, judgment, or ability. Make eye contact with the driver.
   Carefully and deliberately use hand signals so they will not startle or confuse motorists or be mistaken for a flagger's direction before moving into traffic.
- Your movements may startle a motorist and cause an accident, so move deliberately. Do not make sudden movements that might confuse a motorist.

#### 6.0 PROCEDURES

The following procedures address surface and subsurface sampling.

## CAUTION

Each situation must be evaluated individually to determine the applicability and necessity for obtaining a utility clearance ticket/dig permit. Common sense dictates, prior to digging or boring with power equipment, no matter what the depth, or digging by hand in a manner that could damage unprotected underground utilities, that a dig permit is required. See SOP HS-1.0, Utility Locating and Excavation Clearance, for additional clarification. If you do not know or are unsure as to whether a ticket is necessary – **Get**the **Ticket**.

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# 6.1 <u>Overview</u>

Soil sampling is an important adjunct to groundwater monitoring. Sampling of the soil horizons above the groundwater table can detect contaminants before they migrate to the water table, and can establish the amount of contamination absorbed or adsorbed on aquifer solids that have the potential of contributing to groundwater contamination.

Soil types can vary considerably on a hazardous waste site. These variations, along with vegetation, can affect the rate of contaminant migration through the soil. It is important, therefore, that a detailed record be maintained during sampling operations, particularly noting sampling locations, depths, and such characteristics as grain size, color, and odor. Subsurface conditions are often stable on a daily basis and may demonstrate only slight seasonal variation especially with respect to temperature, available oxygen and light penetration. Changes in any of these conditions can radically alter the rate of chemical reactions or the associated microbiological community, thus further altering specific site conditions. Certain vegetation species can create degradation products that can alter contaminant concentrations in soil. This is why vegetation types and extent of degradation of this foliage must be recorded. To prevent degradation, samples must be kept at their at-depth temperature or lower, protected from direct light, sealed tightly in approved glass containers, and be analyzed as soon as possible after collection. In addition, to the extent possible, vegetation should be removed from the sample.

The physical properties of the soil, its grain size, cohesiveness, associated moisture, and such factors as depth to bedrock and water table, will limit the depth from which samples can be collected and the method required to collect them. It is the intent of this document to present the most commonly employed soil sampling methods used at hazardous waste sites.

#### 6.2 Soil Sample Collection

# 6.2.1 Procedure for Preserving and Collecting Soil Samples for Volatile Organic Compound Analysis

Samples collected using traditional methods such as collection in a jar with no preservation have been known to yield non-representative samples due to loss of volatile organic compounds (VOCs). To prevent such losses, preservation of samples with methanol or sodium bisulfate may be used to minimize volatilization and biodegradation. This preservation may be performed either in the field or laboratory, depending on the sampling methodology employed. Because of the large number of sampling methods and associated equipment required, careful coordination between field and laboratory personnel is needed.

Soil samples to be preserved by the laboratory are currently being collected using Method SW-846, 5035. For samples preserved in the field, laboratories are currently performing low-level analyses (sodium bisulfate preservation) and high- to medium-level analyses (methanol preservation) depending on the needs of the end user.

The following procedures outline the necessary steps for collecting soil samples to be preserved at the laboratory, and for collecting soil samples to be preserved in the field with methanol or sodium bisulfate.

# 6.2.1.1 Soil Samples to be Preserved at the Laboratory

Soil samples collected for volatile organic analysis that are to be preserved at the laboratory shall be obtained using a hermetically sealed sample vial such as an EnCore™ sampler. Each sample shall be

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obtained using a reusable sampling handle (T-handle) that can be provided with the EnCore™ sampler when requested and purchased. Collect the sample in the following manner for each EnCore™ sampler:

- 1. Scene Safety Evaluate the area where sampling will occur. Ensure that the area is safe from physical, chemical, and natural hazards. Clear or barricade those hazards that have been identified.
- 2. Wear the appropriate personal protective equipment (PPE). This will include, at a minimum, safety glasses and nitrile surgeon's gloves. If you must kneel on the ground or place equipment on the surface being sampled, cover the ground surface with plastic to minimize surface contamination of your equipment and clothing. Wear knee pads to protect your knees from kneeling on hard or uneven surfaces.
- 3. Load the Encore™ sampler into the T-handle with the plunger fully depressed.
- 4. Expose the area to be sampled using a hand trowel or similar device to remove surface debris.
- 5. Press the T-handle against the freshly exposed soil surface, forcing soil into the sampler. The plunger will be forced upward as the cavity fills with soil.
- 6. When the sampler is full, rotate the plunger and lock it into place. If the plunger does not lock, the sampler is not full. This method ensures there is no headspace. Soft soil may require several plunges or forcing soil against a hard surface such as a sample trowel to ensure that headspace is eliminated.
- 7. Use a paper towel to remove soil from the side of the sampler so a tight seal can be made between the sample cap and the rubber O-ring.
- 8. With soil slightly piled above the rim of the sampler, force the cap on until the catches hook the side of the sampler.
- 9. Remove any surface soil from the outside of the sampler and place in the foil bag provided with the sampler. Good work hygiene practices and diligent decontamination procedures prevents the spread of contamination even on the outside of the containers.
- 10. Label the bag with appropriate information in accordance with SOP SA-6.3.
- 11. Place the full sampler inside a lined cooler with ice and cool to 4°C ± 2°C. Make sure any required trip blanks and temperature blanks are also in the cooler. Secure custody of the cooler in accordance with SOP SA-6.3.
- 12. Typically, collect three Encore™ samplers at each location. Consult the SAP or laboratory to determine the required number of Encore™ samplers to be collected.
- 13. The T-handle shall be decontaminated before moving to the next interval or location using a soap and water wash and rinse, and where applicable, the selected solvent as defined in the project planning documents.

Using this type of sampling device eliminates the need for field preservation and the shipping restrictions associated with preservatives. A complete set of instructions is included with each Encore™ sampler.

After the Encore™samples are collected, they should be placed on ice immediately and delivered to the laboratory within 48 hours (following the chain-of-custody and documentation procedures outlined in SOP SA-6.1). Samples must be preserved by the laboratory within 48 hours of sample collection.

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# 6.2.1.2 <u>Soil Samples to be Preserved in the Field</u>

Soil samples preserved in the field may be prepared for analyses using both the low-level (sodium bisulfate preservation) and high- to medium-level (methanol preservation) methods.

## Safety Reminder

When using chemicals in the field to preserve samples, the FOL and/or SSO must ensure that Materials Safety Data Sheets (MSDSs) have been provided with the chemicals to be used. They also must ensure that these chemicals have been added to the Chemical Inventory List contained within Section 5.0, Hazard Communication, of your Health and Safety Guidance Manual (HSGM). Lastly, but most importantly, the FOL and/or SSO must review the hazards with personnel using these chemicals and ensure that provisions are available for recommended PPE and emergency measures (e.g., eyewash, etc.).

## **Methanol Preservation (High to Medium Level):**

Bottles may be pre-spiked with methanol in the laboratory or prepared in the field. Soil samples to be preserved in the field with methanol shall utilize 40 to 60 mL glass vials with septum-lined lids. Each sample bottle shall be filled with 25 mL of demonstrated analyte-free purge-and-trap grade methanol. The preferred method for adding methanol to the sample bottle is by removing the lid and using a pipette or scaled syringe to add the methanol directly to the bottle.

## **CAUTION**

NEVER attempt to pipette by mouth

In situations where personnel are required to spike the septum using a hypodermic needle, the following provisions for handling sharps must be in place:

- Training of personnel regarding methods for handling of sharps
- Hard-sided containers for the disposal of sharps
- Provisions for treatment in cases where persons have received a puncture wound

Soil shall be collected with the use of a decontaminated (or disposable), small-diameter coring device such as a disposable tube/plunger-type syringe with the tip cut off. The outside diameter of the coring device must be smaller than the inside diameter of the sample bottle neck.

A small electronic balance or manual scale will be necessary for measuring the volume of soil to be added to the methanol-preserved sample bottle. Calibration of the scale shall be performed prior to use and intermittently throughout the day according to the manufacturer's requirements.

The sample should be collected as follows:

- 1. Weigh the unused syringe and plunger to the nearest 0.01 gram.
- 2. Pull the plunger back and insert the syringe into the soil to be sampled.
- 3. Collect 8 to 12 grams of soil by pushing the syringe barrel into the soil.
- 4. Weigh the sample and adjust until obtaining the required amount of sample.

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- Record the sample weight to the nearest 0.01 gram in the field logbook and/or on the sample log sheet.
- 6. Extrude the weighed soil sample into the methanol-preserved sample bottle taking care not to contact the sample container with the syringe.
- 7. If dirty, wipe soil particles from the threads of the bottle and cap. Cap the bottle tightly.
- 8. After capping the bottle, swirl the sample (do not shake) in the methanol and break up the soil such that all of the soil is covered with methanol.
- 9. Place the sample on ice immediately and prepare for shipment to the laboratory as described in SOP SA-6.1.

# **Sodium Bisulfate Preservation (Low Level):**

#### **CAUTION**

Care should be taken when adding the soil to the sodium bisulfate solution. A chemical reaction of soil containing carbonates (limestone) may cause the sample to effervesce or the vial to possibly explode. To avoid this hazard or hazards of this type, a small sample aliquot should be subjected to the sample preservative. If it effervesces in an open air environment, utilize an alternative method such as Encore™ or 2-ounce jar.

Bottles may be prepared in the laboratory or in the field with sodium bisulfate solution. Samples to be preserved in the field using the sodium bisulfate method are to be prepared and collected as follows:

- 1. Add 1 gram of sodium bisulfate to 5 mL of laboratory-grade deionized water in a 40 to 60 mL glass vial with septum-lined lid.
- 2. Collect the soil sample and record the sample weight to the nearest 0.01 gram in the field logbook or on the sample log sheet as described for methanol preservation
- 3. Add the weighed sample to the sample vial.
- 4. Collect duplicate samples using the methanol preservation method on a one-for-one sample basis because it is necessary for the laboratory to perform both low-level and medium-level analyses.
- 5. Place the samples on ice immediately and prepare for shipment to the laboratory as described in SOP SA-6.1.

#### NOTE

If lower detection limits are necessary, an option to field preserving with sodium bisulfate may be to collect EnCore™ samplers at a given sample location. Consult the planning documents to determine whether this is required. If it is, collect samples in accordance with the Encore™ sampling procedure above and then send all samplers to the laboratory to perform the required preservation and analyses.

## 6.2.2 Procedure for Collecting Soil Samples for Non-Volatile Analyses

Samples collected for non-volatile analyses may be collected as either grab or composite samples as follows:

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- 1. With a stainless steel trowel or other approved tool, transfer a portion of soil to be sampled to a stainless steel bowl or disposable inert plastic tray.
- 2. Remove roots, vegetation, sticks, and stones larger than the size of a green pea.
- 3. Thoroughly mix the soil in the bowl or tray to obtain as uniform a texture and color as practicable. The soil type, moisture content, amount of vegetation, and other factors may affect the amount of time required to obtain a properly mixed sample. In some cases, it may be impossible to obtain a uniform sample appearance. Use the field logbook to describe any significant difficulties encountered in obtaining a uniform mixture.
- 4. Transfer the mixed soil to the appropriate sample containers and close the containers.
- 5. Label the sample containers in accordance with SOP SA-6.3.
- 6. Place the containers in a cooler of ice as soon after collection as possible.
- 7. Prepare the sample shipment and ship the samples in accordance with SOP SA-6.1.

## **NOTE**

Cooling may not be required for some samples depending on the scheduled analyses.

Consult the planning documents if in doubt regarding correct sample preservation conditions. When in doubt – Cool to 4°C.

# NOTE

Head space is permitted in soil sample containers for non-volatile analyses to allow for sample expansion.

## 6.2.3 Procedure for Collecting Undisturbed Soil Samples

## NOTE

Use of thin-walled undisturbed tube samplers is restricted by the consistency of the soil to be sampled. Often, very loose and/or wet samples cannot be retrieved by the samplers, and soil with a consistency in excess of very stiff cannot be penetrated by the sampler. Devices such as Dennison or Pitcher core samplers can be used to obtain undisturbed samples of stiff soil. Using these devices normally increases sampling costs, and therefore their use should be weighed against the need for acquiring an undisturbed sample. These devices are not discussed in this SOP because they are not commonly used.

When it is necessary to acquire undisturbed samples of soil for purposes of engineering parameter analysis (e.g., permeability), a thin-walled, seamless tube sampler (Shelby tube) shall be employed using the following collection procedure:

- 1. In preparation for sampling utilizing a drill rig, field personnel must complete the following activities:
  - Ensure that all subsurface drilling activities are preceded by a utility clearance for the area to be investigated. This includes activities described in SOP HS-1.0, Utility Location and Excavation Clearance, as well as any location-specific procedures that may apply.

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# REMEMBER

If you are digging near a marked utility (within the diameter of an underground utility that has been marked plus 18 inches), you must first locate the utility through vacuum extraction or hand digging to ensure that your activities will not damage the utility.

- Complete an Equipment Inspection Checklist for the drill rig or direct-push technology (DPT) rig. This checklist will be provided in the HASP.
- Review the Safe Work Permit prior to conducting the activity.
- Review the activity to be conducted.
- 2. Remove all surface debris (e.g., vegetation, roots, twigs, etc.) from the specific sampling location and drill and/or clean out the borehole to the desired sampling depth. Be careful to minimize potential disturbance of the material to be sampled. In saturated material, withdraw the drill bit slowly to prevent loosening of the soil around the borehole and to maintain the water level in the hole at or above groundwater level.

## **CAUTION**

The use of bottom-discharge bits or jetting through an open-tube sampler to clean out the borehole shall not be allowed. Only the use of side-discharge bits is permitted.

- 3. Determine whether a stationary piston-type sampler is required to limit sample disturbance and aid in retaining the sample. Either the hydraulically operated or control rod activated-type of stationary piston sampler may be used.
- 4. Prior to inserting the tube sampler into the borehole, check to ensure that the sampler head contains a check valve. The check valve is necessary to keep water in the rods from pushing the sample out the tube sampler during sample withdrawal. In addition, the check valve maintains a positive suction within the tube to help retain the sample.
- 5. A stainless steel tube sampler is typically used to minimize chemical reaction between the sample and the sampling tube.
- 6. With the sampling tube resting on the bottom of the hole and the water level in the boring at groundwater level or above, push the tube into the soil with a continuous and rapid motion, without impacting or twisting. If the soil is too hard to penetrate by pushing alone, careful hammering may be used by minimizing drop distance (tapping) of the hammer. Before pulling the tube, turn it at least one revolution to shear the sample off at the bottom. In no case shall the tube be pushed farther than the length provided for the soil sample. Allow about 3 inches in the tube for cuttings and sludge.
- 7. Upon removal of the sampling tube from the hole, measure the length of sample in the tube and also the length penetrated.
- 8. Remove disturbed material in the upper end of the tube and measure the length of sample again.
- 9. After removing at least 1 inch of soil from the lower end, place enough packing material (clean inert material such as paper or cloth) tightly in each end of the Shelby tube and then pour melted wax into each end to make at least a ½-inch wax plug and then add more packing material to fill the voids at both ends.

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- 10. Place plastic caps on the ends, tape the caps in place, and dip the ends in wax to prevent loss of soil.
- 11. Affix label(s) to the tube as required and record sample number, depth, penetration, and recovery length on the label.
- 12. Mark the "up" direction on the side and upper end of the tube with indelible ink.
- 13. Complete a chain-of-custody form (see SOP SA-6.3) and other required forms (including Attachment A of this SOP).
- 14. Ship samples protected with suitable resilient packing material to reduce shock, vibration, and disturbance.

#### **CAUTION**

To preserve sample integrity do not allow tubes to freeze, and store the samples vertically with the same orientation they had in the ground, (i.e., top of sample is up) in a cool place out of the sun at all times.

## CAUTION

A primary concern in the preparation of the wax plugs is the potential for the heat source and melted wax to cause a fire and/or burns. Follow the directions below to prevent injury or fire.

## **Electrical Heating**

Using hot plates to melt the wax is acceptable. In an outdoor setting, make sure a Ground Fault Circuit Interrupter (GFCI) is employed within the electrical circuit. If a portable generator is used, ensure that the generator is an adequate distance from the sampling operation (at least 50 feet). Ensure that the extension cord is rated for the intended load and for outdoor use and is free from recognizable damage. Ensure flammable preservatives are not employed or stored near the hot plate. Although a Hot Work Permit is not required, scene safety evaluation by site personnel of the above elements is. As always, if a fire potential exists, the provisions for extinguishing must be immediately accessible as well as any provisions for first aid measures.

## Open Flame

If an open flame is used, the following provisions are necessary:

- Complete a Hot Work Permit and any local permit required for elevated temperature applications. The Hot Work Permit, provided in your HASP, will aid the FOL and/or the SSO in ensuring that fire protection provisions (extinguishers, fire watches, etc.) are in place as well as ensuring that local requirements have been addressed.
- Ensure that water is available to address any wax splashes or contact. If possible, immerse the contacted area. Where this is not possible, run water over the area and apply cold compresses. The need for medical attention or first aid shall be determined on site under the direction of the SSO.

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# 6.3 Surface Soil Sampling

The simplest, most direct method of collecting surface soil samples for subsequent analysis is by use of a stainless steel shovel, hand auger, soil corer, or stainless steel or disposable plastic trowel.

### NOTE

Multiple depth intervals are used to describe surface soil. Sometimes surface soil is defined as soil from 0 to 2 inches below ground surface (bgs), and sometimes it is defined as soil from other depths such as 0 to 2 feet bgs. Ensure that the definition of surface soil depth is clear before collecting surface soil samples.

For the purposes of instruction, the terms "surface soil" and "near-surface soil" are used in this SOP as follows:

- Surface soil 0 to 6 inches bgs
- Near-surface soil 6 to 18 inches bgs

If these intervals are defined differently in the planning documents, substitute the appropriate depth ranges.

In general, the following equipment is necessary for obtaining surface soil samples:

- Stainless steel or pre-cleaned disposable trowel.
- Stainless steel hand auger, soil corer, or shovel.
- Real-time air monitoring instrument (e.g., PID, FID) as directed in project planning document.
- Required PPE.
  - Nitrile surgeon's or latex gloves may be used, layered as necessary.
  - Safety glasses
  - Other Items identified on the Safe Work Permit may be required based on location-specific requirements such as hearing protection, steel-toed work boots, and a hard hat when working near a drill rig. These provisions will be listed in the HASP or directed by the FOL and/or SSO.

## Safety Reminder

The use of latex products may elicit an allergic reaction in some people. Should this occur, remove the latex gloves, treat for an allergic reaction, and seek medical attention as necessary.

- Required paperwork (see SOP SA-6.3 and Attachment A of this SOP)
- Required decontamination equipment
- Required sample container(s)
- Wooden stakes or pin flags

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- Sealable polyethylene bags (e.g., Ziploc<sup>®</sup> baggies)
- Heavy duty cooler
- Ice
- Chain-of-custody records and custody seals

When acquiring surface soil samples, use the following procedure:

- 1. Place padding or use knee pads when kneeling near the sample location. If necessary, place plastic sheeting to provide a clean surface for sample equipment to avoid possible cross- contamination.
- 2. Carefully remove vegetation, roots, twigs, litter, etc. to expose an adequate soil surface area to accommodate sample volume requirements.
- 3. Using a precleaned syringe or EnCore<sup>TM</sup> samplers, follow the procedure in Section 6.2.1 for collecting surface soil samples for volatile analysis. Surface soil samples for volatile organic analysis should be collected deeper than 6 inches bgs because shallower material has usually lost most of the volatiles through evaporation. Ensure that the appropriate surface soil depth is being analyzed in accordance with the planning document.
- 4. Using decontaminated sampling tools, thoroughly mix in place a sufficient amount of soil to fill the remaining sample containers. See Section 6.5 of this procedure for hand auger instruction, as needed.
- 5. Transfer the sample into those containers utilizing a stainless steel trowel.
- 6. Cap and securely tighten all sample containers.
- 7. Affix a sample label to each container. Be sure to fill out each label carefully and clearly, addressing all the categories described in SOP SA-6.3.
- 8. Proceed with the handling and processing of each sample container as described in SOP SA-6.2.
- 9. Site restoration Whenever removing sample materials, always restore the surface. It is our intent to leave the area better than we found it. Do NOT create trip hazards in areas when pedestrian traffic may exist.

# 6.4 Near-Surface Soil Sampling

Collection of samples from near the surface (depth of 6 to 18 inches) can be accomplished with tools such as shovels, hand auger, soil corers, and stainless steel or pre-cleaned disposable trowels and the equipment listed under Section 6.5 of this procedure.

To obtain near-surface soil samples, the following protocol shall be used:

- 1. With a clean shovel, make a series of vertical cuts in the soil to the depth required to form a square approximately 1 foot by 1 foot.
- 2. Lever out the formed plug and scrape the bottom of the freshly dug hole with a decontaminated stainless steel or pre-cleaned disposable trowel to remove any loose soil.

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3. Follow steps 1 through 9 of Section 6.3.

## 6.5 Subsurface Soil Sampling With a Hand Auger

A hand augering system generally consists of a variety of stainless steel bucket bits (approximately 6.5 inches long and 2, 2.75, 3.25, and 4 inches in diameter), series of extension rods (available in 2-, 3-, 4- and 5-inch lengths), and a T-handle connected to extension rods and to the auger bucket. A larger-diameter bucket bit is commonly used to bore a hole to the desired sampling depth and then it is withdrawn. The larger-diameter bit is then replaced with a smaller-diameter bit, lowered down the hole, and slowly turned into the soil to the completion depth (approximately 6 inches). The apparatus is then withdrawn and the soil sample collected.

The hand auger can be used in a wide variety of soil conditions. It can be used to sample soil either from the surface, or to depths in excess of 12 feet. However, the presence of subsurface rocks and landfill material and collapse of the borehole normally limit sampling depth.

To accomplish soil sampling using a hand augering system, the following equipment is required:

- Complete hand auger assembly (variety of bucket bit sizes)
- Stainless steel mixing bowls
- The equipment listed in Section 6.3
- Miscellaneous hand tools as required to assemble and disassemble the hand auger units

## **CAUTION**

Potential hazards associated with hand augering include:

- Muscle strain and sprain due to over twisting and/or over compromising yourself.
- Equipment failure due to excessive stress on the T-handle or rods through twisting. Failure of any of these components will result in a sudden release and potential injury due to that failure.

As in all situations, any intrusive activities that could damage underground utilities shall be proceeded by a Dig/Excavation permit/ticket. Call the Utility Locating service in the area or your Project Health and Safety Officer for more information. When in doubt – **Get the Ticket!** 

To obtain soil samples using a hand auger, use the following procedure:

- 1. Wearing designated PPE, attach a properly decontaminated bucket bit to a clean extension rod and attach the T-handle to the extension rod.
- 2. Clear the area to be sampled of any surface debris (vegetation, twigs, rocks, litter, etc.).
- 3. Twist the bucket into the ground while pushing vertically downward on the auger. The cutting shoes fill the bucket as it is advanced into the ground.
- 4. As the auger bucket fills with soil, periodically remove any unneeded soil.

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- 5. Add rod extensions as necessary to extend the reach of the auger. Also, note (in a field notebook, boring log, and/or on a standardized data sheet) any changes in the color, texture or odor of the soil as a function of depth. The project-specific planning document (SAP, HASP, etc.) describe requirements for scanning the soil with a real-time air monitoring instrument (e.g., PID, FID, etc.) and recording the measurements.
- 6. After reaching the desired depth (e.g., the top of the interval to be sampled), slowly and carefully withdraw the apparatus from the borehole to prevent or minimize movement of soil from shallower intervals to the bottom of the hole.
- 7. Remove the soiled bucket bit from the rod extension and replace it with another properly decontaminated bucket bit. The bucket bit used for sampling is to be smaller in diameter than the bucket bit employed to initiate the borehole.
- 8. Carefully lower the apparatus down the borehole. Care must be taken to avoid scraping the borehole sides.
- 9. Slowly turn the apparatus until the bucket bit is advanced approximately 6 inches.
- 10. Discard the top of the core (approximately 1 inch), which represents any loose material collected by the bucket bit before penetrating the sample material.
- 11. Using a precleaned syringe or EnCore<sup>™</sup> samplers, follow the procedure in Section 6.2.1 for collecting a soil sample for volatile compound analysis directly from the bucket bit.
- 12. Utilizing a properly decontaminated stainless steel trowel or dedicated disposable trowel, remove the remaining sample material from the bucket bit and place into a properly decontaminated stainless steel mixing bowl.
- 13. Homogenize the sample material as thoroughly as practicable then fill the remaining sample containers. Refer to Section 6.2.2.
- 14. Follow steps 4 through 7 listed in Section 6.3.

## 6.5.1 Sampling Using Stainless Steel Soil Corers

A soil corer is a stainless steel tube equipped with a cutting shoe and sample window in the side. The soil corer is advanced into the soil by applying downward pressure (body weight). The soil is unloaded by then forcing a ram towards the cutting shoe, which results in the discharge of the soil core through a window in the sleeve.

Use, application, and sample protocol is the same as for hand augering provided above, but without necessarily rotating the corer while advancing it.

#### **SAFETY REMINDER**

Hand augering and soil corer sampling can be physically demanding based on the type of geology and subsurface encumbrances encountered. Soil coring has some added hazards such the corer collapsing under your weight. To reduce the potential for muscle strain and damage, the following measures will be incorporated:

- Stretch and limber your muscles before heavy exertion. This hazard becomes more predominant in the early morning hours (prior to muscles becoming limber) and later in the day (as a result of fatigue).

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- Job rotation Share the duties so that repetitive actions do not result in fatigue and injury.
- Increase break frequencies as needed, especially as ambient conditions of heat and/or cold stress may dictate.
- Do not force the hand tools or use cheater pipes or similar devices to bypass an obstruction. Move to another location near the sampling point. Exerting additional forces on the sampling devices can result in damage and/or failure that could potentially injure someone in the immediate vicinity.
- Do not over compromise yourself when applying force to the soil corer or hand auger. If there is a sudden release, it could result in a fall or muscle injury due to strain.

# 6.6 <u>Subsurface Soil Sampling with a Split-Barrel Sampler</u>

A split-barrel (split-spoon) sampler consists of a heavy carbon steel or stainless steel sampling tube that can be split into two equal halves to reveal the soil sample (see Attachment B). A drive head is attached to the upper end of the tube and serves as a point of attachment for the drill rod. A removable tapered nosepiece/drive shoe attaches to the lower end of the tube and facilitates cutting. A basket-like sample retainer can be fitted to the lower end of the split tube to hold loose, dry soil samples in the tube when the sampler is removed from the drill hole. This split-barrel sampler is made to be attached to a drill rod and forced into the ground by means of a 140-pound or larger casing driver.

# Safety Reminder

It is intended through the Equipment Inspection for Drill Rigs form provided in the HASP that the hammer and hemp rope, where applicable, associated with this activity will be inspected (no physical damage is obvious), properly attached to the hammer (suitable knots or sufficient mechanical devices), and is in overall good condition.

Split-barrel samplers are used to collect soil samples from a wide variety of soil types and from depths greater than those attainable with other soil sampling equipment.

The following equipment is used for obtaining split-barrel samples:

- Drilling equipment (provided by subcontractor).
- Split-barrel samplers (2-inch OD, 1-3/8-inch ID, either 20 inches or 26 inches long); Larger OD samplers are available if a larger volume of sample is needed.
- Drive weight assembly, 140-pound weight, driving head, and guide permitting free fall of 30 inches.
- Stainless steel mixing bowls.
- Equipment listed in Section 6.3.

The following steps shall be followed to obtain split-barrel samples (Steps 1 through 4 are typically performed by the drilling subcontractor):

1. Attach the split-barrel sampler to the sampling rods.

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- 2. Lower the sampler into the borehole inside the hollow stem auger bits.
- 3. Advance the split-barrel sampler by hammering the length (typically 18 or 24 inches) of the split-barrel sampler into the soil using 140-pound or larger hammer.
- 4. When the desired depth is achieved, extract the drill rods and sampler from the augers and/or borehole.
- 5. Detach the sampler from the drill rods.
- 6. Place the sampler securely in a vise so it can be opened using pipe wrenches.

#### **CAUTION**

Pipe wrenches are used to separate the split spoon into several components. The driller's helper should not apply excessive force through the use of cheater pipes or push or pull in the direction where, if the wrench slips, hands or fingers will be trapped against an immovable object.

- 7. Remove the drive head and nosepiece with the wrenches, and open the sampler to reveal the soil sample.
- 8. Immediately scan the sample core with a real-time air monitoring instrument (e.g., FID, PID, etc.) (as project-specific planning documents dictate). Carefully separate (or cut) the soil core, with a decontaminated stainless steel knife or trowel, at about 6-inch intervals while scanning the center of the core for elevated readings. Also scan stained soil, soil lenses, and anomalies (if present), and record readings.
- 9. If elevated vapor readings were observed, collect the sample scheduled for volatile analysis from the center of the core where elevated readings occurred. If no elevated readings where encountered, the sample material should be collected from the core's center (this area represents the least disturbed area with minimal atmospheric contact) (refer to Section 6.2.1).
- 10. Using the same trowel, remove remaining sample material from the split-barrel sampler (except for the small portion of disturbed soil usually found at the top of the core sample) and place the soil into a decontaminated stainless steel mixing bowl.
- 11. Homogenize the sample material as thoroughly as practicable then fill the remaining sample containers (refer to Section 6.2.2).
- 12. Follow steps 4 through 7 in Section 6.3.

### 6.7 Subsurface Soil Sampling Using Direct-Push Technology

Subsurface soil samples can be collected to depths of 40+ feet using DPT. DPT equipment, responsibilities, and procedures are described in SOP SA-2.5.

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# 6.8 <u>Excavation and Sampling of Test Pits and Trenches</u>

# 6.8.1 Applicability

This subsection presents routine test pit or trench excavation techniques and specialized techniques that are applicable under certain conditions.

# **CAUTION**

During the excavation of trenches or pits at hazardous waste sites, several health and safety concerns arise from the method of excavation. No personnel shall enter any test pit or excavation over 4 feet deep except as a last resort, and then only under direct supervision of a Competent Person (as defined in 29 CFR 1929.650 of Subpart P -Excavations). Whenever possible, all required chemical and lithological samples should be collected using the excavator bucket or other remote sampling apparatus. If entrance is required, all test pits or excavations must be stabilized by bracing the pit sides using specifically designed wooden, steel, or aluminum support structures or through sloping and benching. Personnel entering the excavation may be exposed to toxic or explosive gases and oxygen-deficient environments; therefore, monitoring will be conducted by the Competent Person to determine if it is safe to enter. Any entry into a trench greater than 4 feet deep will constitute a Confined Space Entry and must be conducted in conformance with OSHA standard 29 CFR 1910.146. In all cases involving entry, substantial air monitoring, before entry, appropriate respiratory gear and protective clothing determination, and rescue provisions are mandatory. There must be at least three people present at the immediate site before entry by one of the field team members. This minimum number of people will increase based on the potential hazards or complexity of the work to be performed. The reader shall refer to OSHA regulations 29 CFR 1926.650, 29 CFR 1910.120, 29 CFR 1910.134, and 29 CFR 1910.146. Highhazard entries such as this will be supported by members of the Health Sciences Group professionally trained in these activities.

Excavations are generally not practical where a depth of more than about 15 to 20-feet is desired, and they are usually limited to a few feet below the water table. In some cases, a pumping system may be required to control water levels within the pit, providing that pumped water can be adequately stored or disposed. If soil data at depths greater than 15-feet are required, the data are usually obtained through test borings instead of test pits.

In addition, hazardous wastes may be brought to the surface by excavation equipment. This material, whether removed from the site or returned to the subsurface, must be properly handled according to any and all applicable federal, state, and local regulations.

#### 6.8.2 Test Pit and Trench Excavation

Test pits or trench excavations are constructed with the intent that they will provide an open view of subsurface lithology and/or disposal conditions that a boring will not provide. These procedures describe the methods for excavating and logging test pits and trenches installed to determine subsurface soil and rock conditions. Test pit operations shall be logged and documented (see Attachment C).

Test pits and trenches may be excavated by hand or power equipment to permit detailed descriptions of the nature and contamination of the in-situ materials. The size of the excavation will depend primarily on the following:

• The purpose and extent of the exploration

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- The space required for efficient excavation
- The chemicals of concern
- The economics and efficiency of available equipment

Test pits normally have a cross section that is 4 to 10 feet square; test trenches are usually 3 to 6 feet wide and may be extended for any length required to reveal conditions along a specific line. The following table provides guidelines for design consideration based on equipment efficiencies.

Equipment	Typical Widths, in Feet	
Trenching machine	0.25 to 1.0	
Backhoe/Track Hoe	2 to 6	

The lateral limits of excavation of trenches and the position of test pits shall be carefully marked on area base maps. If precise positioning is required to indicate the location of highly hazardous materials, nearby utilities, or dangerous conditions, the limits of the excavation shall be surveyed. Also, if precise determination of the depth of buried materials is needed for design or environmental assessment purposes, the elevation of the ground surface at the test pit or trench location shall also be determined by survey. If the test pit/trench will not be surveyed immediately, it shall be backfilled and its position identified with stakes placed in the ground at the margin of the excavation for later surveying.

The construction of test pits and trenches shall be planned and designed in advance as much as possible. However, the following field conditions may necessitate revisions to the initial plans:

- Subsurface utilities
- Surface and subsurface encumbrances
- Vehicle and pedestrian traffic patterns
- Purpose for excavation (e.g., the excavation of potential ordnance items)

The final depth and construction method shall be collectively determined by the FOL and designated Competent Person. The actual layout of each test pit, temporary staging area, and spoils pile may further be predicated based on site conditions and wind direction at the time the test pit is excavated. Prior to excavation, the area may be surveyed by magnetometer or metal detector or other passive methods specified in SOP HS1.0, Utility Location and Excavation Clearance, to identify the presence of underground utilities or drums. Where possible, the excavator should be positioned upwind and preferably within an enclosed cab.

No personnel shall enter any test pit or excavation except as a last resort, and then only under direct supervision of a Competent Person. If entrance is required, OSHA requirements must be met (e.g., walls must be braced with wooden or steel braces, ladders must be placed for every 25 feet of lateral travel and extended 3 feet above ground surface). A temporary guard rail or vehicle stop must be placed along the surface of the hole before entry in situations where the excavation may be approached by traffic. Spoils will be stockpiled no closer than 2 feet from the sidewall of the excavation. The excavation equipment operator shall be careful not to undercut sidewalls and will, where necessary, bench back to increase stability. The top cover, when considered clean, will be placed separately from the subsurface materials to permit clean cover. It is emphasized that the project data needs should be structured such that required samples can be collected without requiring entrance into the excavation. For example,

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samples of leachate, groundwater, or sidewall soil can be collected with telescoping poles or similar equipment.

Dewatering and watering may be required to ensure the stability of the side walls, to prevent the bottom of the pit from heaving, and to keep the excavation stable. This is an important consideration for excavations in cohesionless material below the groundwater table and for excavations left open greater than a day. Liquids removed as a result of dewatering operations must be handled as potentially contaminated materials. Procedures for the collection and disposal of such materials should be discussed in the site-specific project plans.

Where possible excavations and test pits shall be opened and closed within the same working day. Where this is not possible, the following engineering controls shall be put in place to control access:

- Trench covers/street plates
- Fences encompassing the entire excavation intended to control access
- Warning signs warning personnel of the hazards
- Amber flashing lights to demarcate boundaries of the excavation at night

Excavations left open will have emergency means to exit should someone accidentally enter.

# 6.8.3 Sampling in Test Pits and Trenches

## 6.8.3.1 General

Log test pits and trenches as they are excavated in accordance with the Test Pit Log presented in Attachment C. These records include plan and profile sketches of the test pit/trench showing materials encountered, their depth and distribution in the pit/trench, and sample locations. These records also include safety and sample screening information.

Entry of test pits by personnel is extremely dangerous, shall be avoided unless absolutely necessary, and can occur only after all applicable health and safety and OSHA requirements have been met as stated above. These provisions will be reiterated as appropriate in the project-specific HASP.

The final depth and type of samples obtained from each test pit will be determined at the time the test pit is excavated. Sufficient samples are usually obtained and analyzed to quantify contaminant distribution as a function of depth for each test pit. Additional samples of each waste phase and any fluids encountered in each test pit may also be collected.

In some cases, samples of soil may be extracted from the test pit for reasons other than waste sampling and chemical analysis, for instance, to obtain geotechnical information. Such information includes soil types, stratigraphy, strength, etc., and could therefore entail the collection of disturbed (grab or bulk) or relatively undisturbed (hand-carved or pushed/driven) samples that can be tested for geotechnical properties. The purposes of such explorations are very similar to those of shallow exploratory or test borings, but often test pits offer a faster, more cost-effective method of sampling than installing borings.

# 6.8.3.2 Sampling Equipment

The following equipment is needed for obtaining samples for chemical or geotechnical analysis from test pits and trenches:

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- Backhoe or other excavating machinery.
- Shovels, picks, hand augers, and stainless steel trowels/disposable trowels.
- Sample container bucket with locking lid for large samples; appropriate bottle ware for chemical or geotechnical analysis samples.
- Polyethylene bags for enclosing sample containers; buckets.
- Remote sampler consisting of 10-foot sections of steel conduit (1-inch-diameter), hose clamps, and right angle adapter for conduit (see Attachment D).

# 6.8.3.3 Sampling Methods

The methods discussed in this section refer to test pit sampling from grade level. If test pit entry is required, see Section 6.8.3.4.

- Excavate the trench or pit in several 0.5- to 1.0-foot depth increments. Where soil types support the
  use of a sand bar cutting plate, use of this device is recommended to avoid potentially snagging
  utilities with the excavator teeth. It is recommended that soil probes or similar devices be employed
  where buried items or utilities may be encountered. This permits the trench floor to be probed prior to
  the next cut.
- After each increment:
  - the operator shall wait while the sampler inspects the test pit from grade level
  - the sampler shall probe the next interval where this is considered necessary. Practical depth increments for lithological evaluations may range from 2 to 4 feet i or where lithological changes are noted.
- The backhoe operator, who will have the best view of the test pit, shall immediately cease digging if:
  - Any fluid phase, including groundwater seepage, is encountered in the test pit
  - Any drums, other potential waste containers, obstructions, or utility lines are encountered
  - Distinct changes of material being excavated are encountered

This action is necessary to permit proper sampling of the test pit and to prevent a breach of safety protocol. Depending on the conditions encountered, it may be required to excavate more slowly and carefully with the backhoe.

For obtaining test pit samples from grade level, the following procedure shall be followed:

- Use the backhoe to remove loose material from the excavation walls and floor to the greatest extent possible.
- Secure the walls of the pit, if necessary. (There is seldom any need to enter a pit or trench that would justify the expense of shoring the walls. All observations and samples should be taken from the ground surface.)

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- Samples of the test pit material are to be obtained either directly from the backhoe bucket or from the material after it has been deposited on the ground, as follows:
  - a. The sampler or FOL shall direct the backhoe operator to remove material from the selected depth or location within the test pit/trench.
  - b. The backhoe operator shall bring the bucket over to a designated location on the sidewall a sufficient distance from the pit (at least 5 feet) to allow the sampler to work around the bucket.
  - c. After the bucket has been set on the ground, the backhoe operator shall either disengage the controls or shut the machine down.
  - d. When signaled by the operator that it is safe to do, the sampler will approach the bucket.
  - e. The soil shall be monitored with a photoionization or flame ionization detector (PID or FID) as directed in the project -specific planning documents.
  - f. The sampler shall collect the sample from the center of the bucket or pile in accordance with surface soil sampling procedures of Section 6.3 or 6.4, as applicable. Collecting samples from the center of a pile or bucket eliminates cross-contamination from the bucket or other depth intervals.
- If a composite sample is desired, several depths or locations within the pit/trench will be selected, and the bucket will be filled from each area. It is preferable to send individual sample bottles filled from each bucket to the laboratory for compositing under the more controlled laboratory conditions. However, if compositing in the field is required, each sample container shall be filled from materials that have been transferred into a mixing bucket and homogenized. Note that homogenization/compositing is not applicable for samples to be subjected to volatile organic analysis.

# **CAUTION**

Care must be exercised when using the remote sampler described in the next step because of potential instability of trench walls. In situations where someone must move closer than 2 feet to the excavation edge, a board or platform should be used to displace the sampler's weight to minimize the chance of collapse of the excavation edge. Fall protection should also be employed when working near the edges or trenches greater than 6 feet deep. An immediate means to extract people who have fallen into the trench will be immediately available. These means may include ladders or rope anchor points.

- Using the remote sampler shown in Attachment D, samples can be taken at the desired depth from the sidewall or bottom of the pit as follows:
  - a. Scrape the face of the pit/trench using a long-handled shovel or hoe to remove the smeared zone that has contacted the backhoe bucket.
  - b. Collect the sample directly into the sample jar, by scraping with the jar edge, eliminating the need for sample handling equipment and minimizing the likelihood of cross-contamination.
  - c. Cap the sample jar, remove it from the remote sampler assembly, and package the sample for shipment in accordance with SOP SA-6.3.
- Complete documentation as described in SOP SA-6.3 and Attachment C of this SOP.

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# 6.8.3.4 In-Pit Sampling

Under rare conditions, personnel may be required to enter the test pit/trench. This is necessary only when soil conditions preclude obtaining suitable samples from the backhoe bucket (e.g., excessive mixing of soil or wastes within the test pit/trench) or when samples from relatively small discrete zones within the test pit are required. This approach may also be necessary to sample any seepage occurring at discrete levels or zones in the test pit that are not accessible with remote samplers.

In general, personnel shall sample and log pits and trenches from the ground surface, except as provided for by the following criteria:

- There are no practical alternative means of obtaining such data.
- The SSO and Competent Person determine that such action can be accomplished without breaching site safety protocol. This determination will be based on actual monitoring of the pit/trench after it is dug (including, at a minimum, measurements of oxygen concentration, flammable gases, and toxic compounds, in that order). Action levels will be provided in project-specific planning documents.
- A company-designated Competent Person determines that the pit/trench is stable trough soil
  classification evaluation/inspections or is made stable (by cutting/grading the sidewalls or using
  shoring) prior to entrance of any personnel. OSHA requirements shall be strictly observed.

If these conditions are satisfied, only one person may enter the pit/trench. On potentially hazardous waste sites, this individual shall be dressed in selected PPE as required by the conditions in the pit. He/she shall be affixed to a harness and lifeline and continuously monitored while in the pit.

A second and possible third individual shall be fully dressed in protective clothing including a self-contained breathing device and on standby during all pit entry operations to support self rescue or assisted self rescue. The individual entering the pit shall remain therein for as brief a period as practical, commensurate with performance of his/her work. After removing the smeared zone, samples shall be obtained with a decontaminated trowel or spoon.

# 6.8.3.5 Geotechnical Sampling

In addition to the equipment described in Section 6.8.3.2, the following equipment is needed for geotechnical sampling:

- Soil sampling equipment, similar to that used in shallow drilled boring (i.e., thin-walled tube samplers), that can be pushed or driven into the floor of the test pit.
- Suitable driving (e,g., sledge hammer) or pushing (e.g., backhoe bucket) equipment used to advance the sampler into the soil.
- Knives, spatulas, and other suitable devices for trimming hand-carved samples.
- Suitable containers (bags, jars, tubes, boxes, etc.), labels, wax, etc. for holding and safely transporting collected soil samples.
- Geotechnical equipment (pocket penetrometer, torvane, etc.) for field testing collected soil samples for classification and strength properties.

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Disturbed grab or bulk geotechnical soil samples may be collected for most soil in the same manner as comparable soil samples for chemical analysis. These collected samples may be stored in jars or plastic-lined sacks (larger samples), which will preserve their moisture content. Smaller samples of this type are usually tested for their index properties to aid in soil identification and classification: larger bulk samples are usually required to perform compaction tests.

Relatively undisturbed samples are usually extracted in cohesive soil using thin-walled tube samplers, and such samples are then tested in a geotechnical laboratory for their strength, permeability, and/or compressibility. The techniques for extracting and preserving such samples are similar to those used in performing Shelby tube sampling in borings, except that the sampler is advanced by hand or backhoe, rather than by a drill rig. Also, the sampler may be extracted from the test pit by excavation around the tube when it is difficult to pull it out of the ground. If this excavation requires entry of the test pit, the requirements described in Section 6.8.3.4 shall be followed. The thin-walled tube sampler shall be pushed or driven vertically into the floor or steps excavated in the test pit at the desired sampling elevations. Extracting tube samples horizontally from the walls of the test pit is not appropriate because the sample will not have the correct orientation.

A sledge hammer or backhoe may be used to drive or push the tube into the ground. Place a piece of wood over the top of the sampler or sampling tube to prevent damage during driving/pushing of the sample. Pushing the sampler with a constant thrust is always preferable to driving it with repeated blows, thus minimizing disturbance to the sample. When using a sledge hammer, it is recommended that the sampler be stabilized using a rope/strap wrench or pipe wrench to remove the person's hands holding the sampler from the strike zone. If the sample cannot be extracted by rotating it at least two revolutions (to shear off the sample at the bottom), hook the sampler to the excavator or backhoe and extract. This means an alternative head will be used as a connection point or that multiple choke hitches will be applied to extract the sampler. If this fails and the excavator can dig deeper without potentially impacting subsurface utilities, excavate the sampler. If this fails or if the excavator cannot be used due to subsurface utilities, hand-excavate to remove the soil from around the sides of the sampler. If hand-excavation requires entry into the test pit, the requirements in Section 6.8.3.4 must be followed. Prepare the sample as described in Steps 9 through 13 in Section 6.2.3, and label, pack and transport the sample in the required manner, as described in SOPs SA-6.3 and SA-6.1.

## 6.8.4 Backfilling of Trenches and Test Pits

All test pits and excavations must be either backfilled, covered, or otherwise protected at the end of each day. No excavations shall remain open during non-working hours unless adequately covered or otherwise protected.

Before backfilling, the onsite crew may photograph, if required by the project-specific work plan, all significant features exposed by the test pit and trench and shall include in the photograph a scale to show dimensions. Photographs of test pits shall be marked to include site number, test pit number, depth, description of feature, and date of photograph. In addition, a geologic description of each photograph shall be entered in the site logbook. All photographs shall be indexed and maintained as part of the project file for future reference.

After inspection, backfill material shall be returned to the pit under the direction of the FOL. Backfill should be returned to the trench or test pit in 6-inch to 1-foot lifts and compacted with the bucket. Remote controlled tampers or rollers may be lowered into the trench and operated from top side. This procedure will continue to the grade surface. It is recommended that the trench be tracked or rolled in. During excavation, clean soil from the top 2 feet may have been separated to be used to cover the last segments. Where these materials are not clean, it is recommended that clean fill be used for the top cover.

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If a low-permeability layer is penetrated (resulting in groundwater flow from an upper contaminated flow zone into a lower uncontaminated flow zone), backfill material must represent original conditions or be impermeable. Backfill could consist of a soil-bentonite mix prepared in a proportion specified by the FOL (representing a permeability equal to or less than original conditions). Backfill can be covered by "clean" soil and graded to the original land contour. Revegetation of the disturbed area may also be required.

# 6.9 Records

The appropriate sample log sheet (see Attachment A of this SOP) must be completed by the site geologist/sampler for all samples collected. All soil sampling locations should be documented by tying in the location of two or more nearby permanent landmarks (building, telephone pole, fence, etc.) or obtaining GPS coordinates; and shall be noted on the appropriate sample log sheet, site map, or field notebook. Surveying may also be necessary, depending on the project requirements.

Test pit logs (see Attachment C of this SOP) shall contain a sketch of pit conditions. If the project-specific work plan requires photographs, at least one photograph with a scale for comparison shall be taken of each pit. Included in the photograph shall be a card showing the test pit number. Boreholes, test pits, and trenches shall be logged by the field geologist in accordance with SOP GH-1.5.

Other data to be recorded in the field logbook include the following:

- Name and location of job
- Date of boring and excavation
- Approximate surface elevation
- Total depth of boring and excavation
- Dimensions of pit
- Method of sample acquisition
- Type and size of samples
- Soil and rock descriptions
- Photographs if required
- Groundwater levels
- PID/FID/LEL/O<sub>2</sub> meter readings
- Other pertinent information, such as waste material encountered

In addition, site-specific documentation to be maintained by the SSO and/or Competent Person will be required including:

- Calibration logs
- Excavation inspection checklists

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# • Soil type classification

## 7.0 REFERENCES

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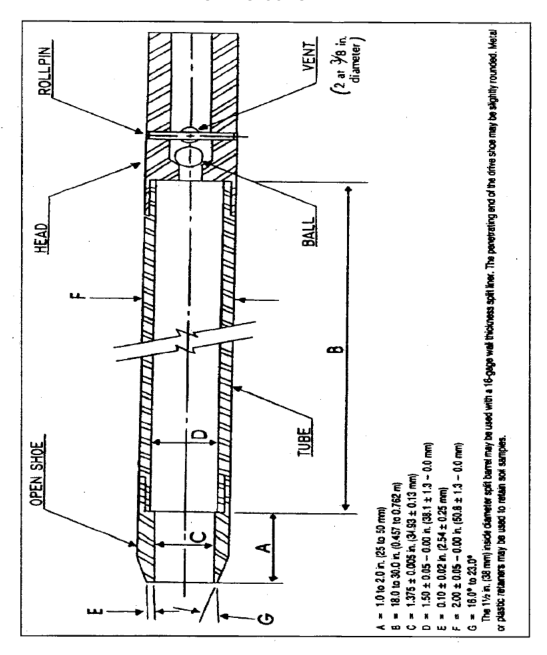
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# ATTACHMENT A SOIL & SEDIMENT SAMPLE LOG SHEET

Tab Tetu	ra Tech NUS,	lnc.	SOIL & SEDII	MENT SAM	PLE LOG SHE	ΈΤ
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Time:						
Method:						
Monitor Reading (pp	m):			1		
COMPOSITE SAMP			- 74 G	1		<b>3</b> 4.
Date:	Time	Depth	Color	Description	n (Sand, Silt, Clay, M	oisture, etc.)
Method:						
Monitor Readings				1		
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(Range in ppm):				<del> </del>		
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SAMPLE COLLECT	ION INFORMATI	ON:				
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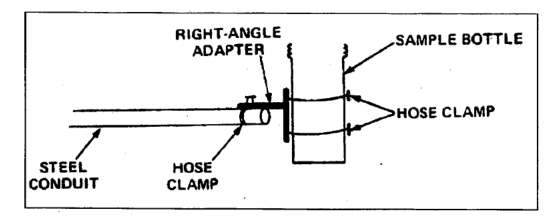
# ATTACHMENT B SPLIT-SPOON SAMPLER



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		ECT NAME:		TE	ST PIT N	o.:		
	LOCA	ECT NUMBER TION:	:	GE	TE: OLOGIS	Г:		
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# ATTACHMENT D REMOTE SAMPLE HOLDER FOR TEST PIT/TRENCH SAMPLING





# STANDARD OPERATING PROCEDURES

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Effective Date 09/03	Revision 3

Applicability

Tetra Tech NUS, Inc.

Prepared

Earth Sciences Department

Approved

D. Senovich

# TETRA TECH NUS, INC.

Subject DIRECT PUSH TECHNOLOGY (GEOPROBE®/HYDROPUNCH™)

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### 1.0 PURPOSE

The purpose of this procedure is to provide general reference information on Direct Push Technology (DPT). DPT is designed to collect soil, groundwater, and soil gas samples without using conventional drilling techniques. The advantage of using DPT over conventional drilling includes the generation of little or no drill cuttings, sampling in locations with difficult accessibility, reduced overhead clearance requirements, no fluid introduction during probing, and typical lower costs per sample than with conventional techniques. Disadvantages include a maximum penetration depth of approximately 15 to 40 feet in dense soils (although it may be as much as 60 to 80 feet in certain types of geological environments), reduced capability of obtaining accurate water-level measurements, and the inability to install permanent groundwater monitoring wells. The methods and equipment described herein are for collection of surface and subsurface soil samples and groundwater samples. Soil gas sampling is discussed in SOP SA-2.4.

#### 2.0 SCOPE

This procedure provides information on proper sampling equipment and techniques for DPT. Review of the information contained herein will facilitate planning of the field sampling effort by describing standard sampling techniques. The techniques described shall be followed whenever applicable, noting that site-specific conditions or project-specific plans may require adjustments in methodology.

### 3.0 GLOSSARY

<u>Direct Push Technology (DPT)</u> - DPT refers to sampling tools and sensors that are driven directly into the ground without the use of conventional drilling equipment. DPT typically utilizes hydraulic pressure and/or percussion hammers to advance the sampling tools. A primary advantage of DPT over conventional drilling techniques is that DPT results in the generation of little or no investigation derived waste.

<u>Geoprobe®</u> - Geoprobe® is a manufacturer of a hydraulically-powered, percussion/probing machines utilizing DPT to collect subsurface environmental samples. Geoprobe® relies on a relatively small amount of static weight (vehicle) combined with percussion as the energy for advancement of a tool string. The Geoprobe® equipment can be mounted in a multitude of vehicles for access to all types of environmental sites.

<u>HydroPunch™</u> - HydroPunch™ is a manufacturer of stainless steel and Teflon® sampling tools that are capable of collecting representative groundwater and/or soil samples without requiring the installation of a groundwater monitoring well or conventional soil boring. HydroPunch™ is an example of DPT sampling equipment.

<u>Flame Ionization Detector (FID)</u> - A portable instrument for the measurement of many combustible organic compounds and a few inorganic compounds in air at parts-per million levels. The basis for the detection is the ionization of gaseous species utilizing a flame as the energizing source.

<u>Photo Ionization Detector (PID)</u> - A portable instrument for the measurement of many combustible organic compounds and a few inorganic compounds in air at parts-per million levels. The basis for the detection is the ionization of gaseous species utilizing ultraviolet radiation as the energizing source.

### 4.0 RESPONSIBILITIES

<u>Project Manager</u> - The Project Manager is responsible for selecting and/or reviewing the appropriate DPT drilling procedure required to support the project objectives.

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<u>Field Operations Leader (FOL)</u>- The FOL is primarily responsible for performing the DPT in accordance with the project-specific plan.

# 5.0 SOIL SAMPLING PROCEDURES

### 5.1 General

The common methodology for the investigation of the vadose zone is soil boring drilling and soil sampling. However, drilling soil borings can be very expensive. Generally the advantage of DPT for subsurface soil sampling is the reduced cost of disposal of drilling cuttings and shorter sampling times.

# 5.2 <u>Sampling Equipment</u>

Equipment needed for conducting DPT drilling for subsurface soil sampling includes, but is not limited to, the following:

- Geoprobe<sup>®</sup> Sampling Kit
- Cut-resistant gloves
- 4-foot x 1.5-inch diameter macrocore sampler
- Probe sampling adapters
- Roto-hammer with 1.5-inch bit
- Disposable acetate liners for soil macrocore sampler
- Cast aluminum or steel drive points
- Geoprobe® AT-660 Series Large Bore Soil Sampler, or equivalent
- Standard decontamination equipment and solutions

For health and safety equipment and procedures, follow the direction provided in the Safe Work Permit in Attachment 1, or the more detailed directions provided in the project's Health and Safety Plan.

## 5.3 DPT Sampling Methodology

There are several methods for the collection of soil samples using DPT drilling. The most common method is discussed in the following section. Variations of the following method may be conducted upon approval of the Project Manager in accordance with the project-specific plan.

- Macrocore samplers fitted with detachable aluminum or steel drive points are driven into the ground using hydraulic pressure. If there is concrete or pavement over a sampling location, a Roto-hammer is used to drill a minimum 1.5-inch diameter hole through the surface material. A Roto-hammer may also be used if very dense soils are encountered.
- The sampler is advanced continuously in 4-foot intervals or less if desired. No soil cuttings are generated because the soil which is not collected in the sampler is displaced within the formation.
- The sampler is retracted from the hole, and the 4-foot continuous sample is removed from the outer coring tube. The sample is contained within an inner acetate liner.
- Attach the metal trough from the Geoprobe® Sampling Kit firmly to the tail gate of a vehicle. If a vehicle with a tail gate is not available, secure the trough on another suitable surface.
- Place the acetate liner containing the soils in the trough.

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- While wearing cut-resistant gloves (constructed of leather or other suitable material), cut the acetate liner through its entire length using the double-bladed knife that accompanies the Geoprobe® Sampling Kit. Then remove the strip of acetate from the trough to gain access to the collected soils. Do not attempt to cut the acetate liner while holding it in your hand.
- Field screen the sample with an FID or PID, and observe/examine the sample (according to SOP GH-1.3). If appropriate, transfer the sample to sample bottles for laboratory analysis. If additional volume is required, push an additional boring adjacent to the first and composite/mix the same interval. Field compositing is usually not acceptable for sample requiring volatile organics analysis.
- Once sampling has been completed, the hole is backfilled with bentonite chips or bentonite cement
  grout, depending upon project requirements. Asphalt or concrete patch is used to cap holes through
  paved or concrete areas. All holes should be finished smooth to existing grade.
- In the event the direct push van/truck cannot be driven to a remote location or a sampling location with difficult accessibility, sampling probes may be advanced and sampled manually or with air/electric operated equipment (e.g., jack hammer).
- Sampling equipment is decontaminated prior to collecting the next sample.

# -6.0 GROUNDWATER SAMPLING PROCEDURES

#### 6.1 General

The most common methodology for the investigation of groundwater is the installation and sampling of permanent monitoring wells. If only groundwater screening is required, the installation and sampling of temporary well points may be performed. The advantage of temporary well point installation using DPT is reduced cost due to no or minimal disposal of drilling cuttings and well construction materials, and shorter installation/times campling.

Two disadvantages of DPT drilling for well point installation are:

- In-aquifors with low yields, well-points may have to be sampled without purging or development.
- If volume requirements are high, this method can be time consuming for low yield aquifers.

### 6.2 <u>Sampling Equipment</u>

Equipment needed for temporary well installation and sampling using DPT includes, but is not limited, to-the following:—

- 2-foot x-1-inch diameter mill-slotted (0.005 to 0.02-inch) well point -
- Connecting rods—
- Rete-hammer with 1.5-inch bit-
- Mechanical jack-
- 1/4-inch OD polyethylene tubing
- 3/8-inch OD polyethylene tubing-
- Peristaltic pump
- Standard decontamination equipment and solutions

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# 6.3 <u>DPT Temporary Well Point Installation and Sampling Methodology</u>

There are several methods for the installation and sampling of temporary well points using DPT. The most common methodology is discussed below. Variations of the following method may be conducted upon approval of the Project Manager in accordance with the project specific plan.

- A 2-foot x 1-inch diameter mill-slotted (0.005 to 0.02-inch) well-point attached to connecting rods isdriven into the ground to the desired depth using a rotary electric hammer or other direct push drill rightlif there is concrete or pavement over a sampling location, a Roto-hammer or electric coring machine is used to drill a hole through the surface material.
- The well-point will be allowed to equilibrate for at least 15 minutes, after which a measurement of the
   -static water level will be taken. The initial measurement of the water level will be used to assess the
   -amount of water which is present in the well-point and to determine the amount of silt-and sand
   infiltration that may have occurred.
- The-well-point-will be developed using a peristaltic pump and polyethylene tubing to remove silt and sand which may have entered the well-point. The well-point is developed by inserting polyethylene-tubing to the bottom of the well-point and lifting and lowering the tubing slightly while the pump isoperating. The pump will be operated at a maximum rate of approximately 2 liters per minute. After removal of sediment from the bottom of the well-point, the well-point will be vigorously pumped at maximum capacity until discharge water is visibly clear and no further sediments are being generated. Measurements of pH, specific conductance, temperature, and turbidity shall be recorded every 5 to 10 minutes during the purging process. After two consistent readings of pH, specific conductance, temperature and turbidity (±10 percent), the well-may be sampled.
- A sample will be collected using the peristaltic pump set at the same or reduced speed as during well development. Samples (with the exception of the samples to be analyzed for volatile organic eempounds, VOCs) will be collected directly from the pump discharge. Sample containers for VOCs will be filled by (first shutting off the pump) crimping the discharge end of the sample tubing when filled, removing the inlet end of the sample tubing from the well, suspending the inlet tubing above the vial, and allowing water to fill each vial by gravity flow.
- Once the groundwater sample has been collected, the connecting rods and well point will be removed from the hole with the direct push rig hydraulies. The hole will be backfilled with bentonite chips or bentonite cement grout, depending upon project requirements. Asphalt or concrete patch will be used to cap holes through paved or concrete areas. All holes will be finished smooth to existing grade.
- In the event the direct push van/truck cannot be driven to a remote location or sampling location with
  difficult accessibility, sampling probes may be advanced and sampled manually or with air/electricoperated equipment (e.g., jack hammer).
- Decontaminate the equipment before moving to the next location.

### 7.0 RECORDS

A record of all field procedures, tests, and observations must be recorded in the field logbook, boring logs, and sample log sheets, as needed. Entries should include all pertinent data regarding the investigation. The use of sketches and field landmarks will help to supplement the investigation and evaluation.

Subject Number Page DIRECT PUSH TECHNOLOGY SA-2.5 6 of 6 (GEOPROBE®/HYDROPUNCH™) Revision Effective Date 09/03 3 **ATTACHMENT 1** SAFE WORK PERMIT FOR DPT OPERATIONS Permit No. Time: From SECTION I: General Job Scope I. Work limited to the following (description, area, equipment used): Monitoring well drilling and installation through direct push technology II. Required Monitoring Instruments: III. Field Crew: On-site Inspection conducted \( \subseteq \text{Yes} \) ☐ No Initials of Inspector SECTION II: General Safety Requirements (To be filled in by permit issuer) Protective equipment required Respiratory equipment required Level D \ Level B \ Level A \ \ Full face APR Escape Pack Half face APR SCBA Detailed on Reverse SKA-PAC SAR Bottle Trailer Skid Rig None Level D Minimum Requirements: Sleeved shirt and long pants, safety footwear, and work gloves Safety glasses, hard hats, and hearing protection will be worn when working near or sampling in the vicinity of the DPT rig. Modifications/Exceptions. VI. Chemicals of Concern Action Level(s) Response Measures VII. Additional Safety Equipment/Procedures Hearing Protection (Plugs/Muffs) Yes
Safety belt/harness
Radio Yes
Barricades Yes Hard-hat ...... ⊠ Yes □ No Safety Glasses ..... ⊠ Yes □ No ⊠ No Chemical/splash goggles..... ☐ Yes ☒ No ⊠ No □ No Gloves (Type -☐ Yes ☐ No Work/warming regimen ☐ Yes Modifications/Exceptions: Reflective vests for high traffic areas. Procedure review with permit acceptors Yes NA Yes NA Safety shower/eyewash (Location & Use)..... Emergency alarms ..... Daily tail gate meetings..... Evacuation routes ..... Contractor tools/equipment/PPE inspected ...... Assembly points ..... IX. Site Preparation Utility Clearances obtained for areas of subsurface investigation ☐ Yes ☐ No Physical hazards removed or blockaded ☐ Yes ☐ No Site control boundaries demarcated/signage ☐ Yes ☐ No **Equipment Preparation** Equipment drained/depressurized..... Equipment purged/cleaned..... Isolation checklist completed..... Electrical lockout required/field switch tested ..... Blinds/misalignments/blocks & bleeds in place ..... Hazardous materials on walls/behind liners considered..... If yes, complete permit required or contact Health Sciences, Pittsburgh Office XII. Special instructions, precautions: Permit Issued by:\_\_\_\_\_ Permit Accepted by:



**TETRA TECH NUS, INC.** 

# STANDARD OPERATING PROCEDURES

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	** * '

Applicability

Tetra Tech NUS, Inc.

Prepared

Earth Sciences Department

Approved

D. Senovich

Subject

NON-RADIOLOGICAL SAMPLE HANDLING

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#### 1.0 PURPOSE

The purpose of this Standard Operating Procedure (SOP) is to provide information on sample preservation, packaging, and shipping procedures to be used in handling environmental samples submitted for chemical constituent, biological, or geotechnical analysis. Sample chain-of-custody procedures and other aspects of field documentation are addressed in SOP SA-6.3. Sample identification is addressed in SOP CT-04.

# 2.0 SCOPE

This procedure describes the appropriate containers to be used for samples depending on the analyses to be performed, and the steps necessary to preserve the samples when shipped off site for chemical analysis.

# 3.0 GLOSSARY

<u>Hazardous Material</u> - A substance or material which has been determined by the Secretary of Transportation to be capable of posing an unreasonable risk to health, safety, and property when transported in commerce, and which has been so designated. Under 49 CFR, the term includes hazardous substances, hazardous wastes, marine pollutants, and elevated temperature materials, as well as materials designated as hazardous under the provisions of §172.101 and §172.102 and materials that meet the defining criteria for hazard classes and divisions in Part 173. With slight modifications, IATA has adopted DOT "hazardous materials" as IATA "Dangerous Goods."

Hazardous Waste - Any substance listed in 40 CFR, Subpart D (y261.30 et seq.), or otherwise characterized as ignitable, corrosive, reactive, or toxic (as defined by Toxicity Characteristic Leaching Procedure, TCLP, analysis) as specified under 40 CFR, Subpart C (y261.20 et seq.), that would be subject to manifest requirements specified in 40 CFR 262. Such substances are defined and regulated by EPA.

<u>Marking</u> - A descriptive name, identification number, instructions, cautions, weight, specification or UN marks, or combination thereof required on outer packaging of hazardous materials.

<u>n.o.i</u> - Not otherwise indicated (may be used interchangeably with n.o.s.).

n.o.s. - Not otherwise specified.

<u>Packaging</u> - A receptacle and any other components or materials necessary for compliance with the minimum packaging requirements of 49 CFR 174, including containers (other than freight containers or overpacks), portable tanks, cargo tanks, tank cars, and multi-unit tank-car tanks to perform a containment function in conformance with the minimum packaging requirements of 49 CFR 173.24(a) & (b).

<u>Placard</u> - Color-coded, pictorial sign which depicts the hazard class symbol and name and which is placed on the side of a vehicle transporting certain hazardous materials.

# Common Preservatives:

- Hydrochloric Acid HCl
- Sulfuric Acid H<sub>2</sub>SO<sub>4</sub>
- Nitric Acid HNO<sub>3</sub>
- Sodium Hydroxide NaOH

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## Other Preservatives

- Zinc Acetate
- Sodium Thiosulfate Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub>

Normality (N) - Concentration of a solution expressed as equivalent per liter, an equivalent being the amount of a substance containing 1 gram-atom of replaceable hydrogen or its equivalent.

Reportable Quantity (RQ) - For the purposes of this SOP, means the quantity specified in column 3 of the Appendix to DOT 49 CFR §172.101 for any material identified in column 1 of the appendix. A spill greater than the amount specified must be reported to the National Response Center.

<u>Sample</u> - A sample is physical evidence collected from a facility or the environment, which is representative of conditions at the location and time of collection.

### 4.0 RESPONSIBILITIES

<u>Field Operations Leader</u> - Directly responsible for the bottling, preservation, labeling, packaging, shipping, and custody of samples up to and including release to the shipper.

<u>Field Samplers</u> - Responsible for initiating the Chain-of-Custody Record (per SOP SA-6.3), implementing the packaging and shipping requirements, and maintaining custody of samples until they are relinquished to another custodian or to the shipper.

### 5.0 PROCEDURES

Sample identification, labeling, documentation, and chain-of-custody are addressed by SOP SA-6.3.

# 5.1 Sample Containers

Different types of chemicals react differently with sample containers made of various materials. For example, trace metals adsorb more strongly to glass than to plastic, whereas many organic chemicals may dissolve various types of plastic containers. Attachments A and B show proper containers (as well as other information) per 40 CFR 136. In general, the sample container shall allow approximately 5-10 percent air space ("ullage") to allow for expansion/vaporization if the sample warms during transport. However, for collection of volatile organic compounds, head space shall be omitted. The analytical laboratory will generally provide certified-clean containers for samples to be analyzed for chemical constituents. Shelby tubes or other sample containers are generally provided by the driller for samples requiring geotechnical analysis. Sufficient lead time shall be allowed for a delivery of sample container orders. Therefore, it is critical to use the correct container to maintain the integrity of the sample prior to analysis.

Once opened, the container must be used at once for storage of a particular sample. Unused but opened containers are to be considered contaminated and must be discarded. Because of the potential for introduction of contamination, they cannot be reclosed and saved for later use. Likewise, any unused containers which appear contaminated upon receipt, or which are found to have loose caps or a missing Teflon liner (if required for the container), shall be discarded.

# 5.2 Sample Preservation

Many water and soil samples are unstable and therefore require preservation to prevent changes in either the concentration or the physical condition of the constituent(s) requiring analysis. Although complete and irreversible preservation of samples is not possible, preservation does retard the chemical and biological

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changes that inevitably take place after the sample is collected. Preservation techniques are usually limited to pH control, chemical addition(s), and refrigeration/ freezing (certain biological samples only).

### 5.2.1 Overview

The preservation techniques to be used for various analytes are listed in Attachments A and B. Reagents required for sample preservation will either be added to the sample containers by the laboratory prior to their shipment to the field or be added in the field (in a clean environment). Only high purity reagents shall be used for preservation. In general, aqueous samples of low-concentration organics (or soil samples of low- or medium-concentration organics) are cooled to 4°C. Medium-concentration aqueous samples, high-hazard organic samples, and some gas samples are typically not preserved. Low-concentration aqueous samples for metals are acidified with HNO<sub>3</sub>, whereas medium-concentration and high-hazard aqueous metal samples are not preserved. Low- or medium-concentration soil samples for metals are cooled to 4°C, whereas high-hazard samples are not cooled.

The following subsections describe the procedures for preparing and adding chemical preservatives. Attachments A and B indicate the specific analytes which require these preservatives.

The FOL is responsible for ensuring that an accurate Chemical Inventory is created and maintained for all hazardous chemicals brought to the work site (see Section 5 of the TtNUS Health and Safety Guidance Manual). Furthermore, the FOL must ensure that a corresponding Material Safety Data Sheet (MSDS) is collected for every substance entered on the site Chemical Inventory, and that all persons using/handling/disposing of these substances review the appropriate MSDS for substances they will work with. The Chemical Inventory and the MSDSs must be maintained at each work site in a location and manner where they are readily-accessible to all personnel.

# 5.2.2 Preparation and Addition of Reagents

Addition of the following acids or bases may be specified for sample preservation; these reagents shall be analytical reagent (AR) grade or purer and shall be diluted to the required concentration with deionized water before field sampling commences. To avoid uncontrolled reactions, be sure to Add Acid to water (not vice versa). A dilutions guide is provided below.

Acid/Base	Dilution	Concentration	Estimated Amount Required for Preservation
Hydrochloric Acid (HCI)	1 part concentrated HCI: 1 part double-distilled, deionized water	6N	5-10 mL
Sulfuric Acid (H <sub>2</sub> SO <sub>4</sub> )	1 part concentrated H <sub>2</sub> SO <sub>4</sub> : 1 part double-distilled, deionized water	18N	2 - 5 mL
Nitric Acid (HNO <sub>3</sub> )	Undiluted concentrated HNO <sub>3</sub>	16N	2 - 5 mL
Sodium Hydroxide (NaOH)	400 grams solid NaOH dissolved in 870 mL double-distilled, deionized water; yields 1 liter of solution	10N	2 mL

The amounts required for preservation shown in the above table assumes proper preparation of the preservative and addition of the preservative to one liter of aqueous sample. This assumes that the sample is initially at pH 7, is poorly buffered, and does not contain particulate matter; as these conditions vary, more preservative may be required. Consequently, the final sample pH must be checked using narrow-range pH paper, as described in the generalized procedure detailed below:

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- Pour off 5-10 mL of sample into a dedicated, clean container. Use some of this sample to check the initial sample pH using wide range (0-14) pH paper. Never dip the pH paper into the sample; always apply a drop of sample to the pH paper using a clean stirring rod or pipette.
- Add about one-half of the estimated preservative required to the original sample bottle. Cap and invert gently several times to mix. Check pH (as described above) using medium range pH paper (pH 0-6 or pH 7.5-14, as applicable).
- · Cap sample bottle and seal securely.

Additional considerations are discussed below:

 To test if ascorbic acid must be used to remove oxidizing agents present in the sample before it can be properly preserved, place a drop of sample on KI-starch paper. A blue color indicates the need for ascorbic acid addition.

If required, add a few crystals of ascorbic acid to the sample and retest with the KI-starch paper. Repeat until a drop of sample produces no color on the KI-starch paper. Then add an additional 0.6 grams of ascorbic acid per each liter of sample volume.

Continue with proper base preservation of the sample as described above.

• Samples for sulfide analysis must be treated by the addition of 4 drops (0.2 mL) of 2N zinc acetate solution per 100 ml of sample.

The 2N zinc acetate solution is made by dissolving 220 grams of zinc acetate in 870 mL of double-distilled, deionized water to make 1 liter of solution.

The sample pH is then raised to 9 using the NaOH preservative.

 Sodium thiosulfate must be added to remove residual chlorine from a sample. To test the sample for residual chlorine use a field test kit specially made for this purpose.

If residual chlorine is present, add 0.08 grams of sodium thiosulfate per liter of sample to remove the residual chlorine.

Continue with proper acidification of the sample as described above.

For biological samples, 10% buffered formalin or isopropanol may also be required for preservation. Questions regarding preservation requirements should be resolved through communication with the laboratory before sampling begins.

### 5.3 Field Filtration

At times, field-filtration may be required to provide for the analysis of dissolved chemical constituents. Field-filtration must be performed <u>prior to</u> the preservation of samples as described above. General procedures for field filtration are described below:

• The sample shall be filtered through a non-metallic, 0.45-micron membrane filter, immediately after collection. The filtration system shall consist of dedicated filter canister, dedicated tubing, and a peristaltic pump with pressure or vacuum pumping squeeze action (since the sample is filtered by mechanical peristalsis, the sample travels only through the tubing).

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- To perform filtration, thread the tubing through the peristaltic pump head. Attach the filter canister to the discharge end of the silicon tubing (note flow direction arrow); attach the aqueous sample container to the intake end of the silicon tubing. Turn the peristaltic pump on and perform filtration. Run approximately 100 ml of sample through the filter and discard prior to sample collection.
- Continue by preserving the filtrate (contained in the filter canister), as applicable and generally described above.

# 5.4 Sample Packaging and Shipping

Only employees who have successfully completed the TtNUS "Shipping Hazardous Materials" training course are authorized to package and ship hazardous substances. These trained individuals are responsible for performing shipping duties in accordance with this training.

Samples collected for shipment from a site shall be classified as either <u>environmental</u> or <u>hazardous</u> <u>material samples</u>. Samples from drums containing materials other than Investigative Derived Waste (IDW) and samples obtained from waste piles or bulk storage tanks are generally shipped as hazardous materials. A distinction must be made between the two types of samples in order to:

- Determine appropriate procedures for transportation of samples (if there is any doubt, a sample shall be considered hazardous and shipped accordingly.)
- Protect the health and safety of transport and laboratory personnel receiving the samples (special precautions are used by the shipper and at laboratories when hazardous materials are received.)

Detailed procedures for packaging environmental samples are outlined in the remainder of this section.

### 5.4.1 Environmental Samples

Environmental samples are packaged as follows:

- Place properly identified sample container, with lid securely fastened, in a plastic bag (e.g. Ziploc baggie), and seal the bag.
- Place sample in a cooler constructed of sturdy material which has been lined with a large, plastic bag (e.g. "garbage" bag). Drain plugs on coolers must be taped shut.
- Pack with enough cushioning materials such as bubble wrap (shoulders of bottles must be iced if required) to minimize the possibility of the container breaking.
- If cooling is required (see Attachments A and B), place ice around sample container shoulders, and on top of packing material (minimum of 8 pounds of ice for a medium-size cooler).
- Seal (i.e., tape or tie top in knot) large liner bag.
- The original (top, signed copy) of the COC form shall be placed inside a large Ziploc-type bag and taped inside the lid of the shipping cooler. If multiple coolers are sent but are included on one COC form, the COC form should be sent with the cooler containing the vials for VOC analysis. The COC form should then state how many coolers are included with that shipment.
- Close and seal outside of cooler as described in SOP SA-6.3. Signed custody seals must be used.

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Coolers must be marked as containing "Environmental Samples." The appropriate side of the container must be marked "This End Up" and arrows placed appropriately. No DOT marking or labeling is required; there are no DOT restrictions on mode of transportation.

### 6.0 REFERENCES

American Public Health Association, 1981. <u>Standard Methods for the Examination of Water and Wastewater</u>, 15th Edition. APHA, Washington, D.C.

International Air Transport Association (latest issue). <u>Dangerous Goods Regulations</u>, Montreal, Quebec, Canada.

- U.S. Department of Transportation (latest issue). Hazardous Materials Regulations, 49 CFR 171-177.
- U.S. EPA, 1984. "Guidelines Establishing Test Procedures for the Analysis of Pollutants under Clean Water Act." Federal Register, Volume 49 (209), October 26, 1984, p. 43234.
- U.S. EPA, 1979. Methods for Chemical Analysis of Water and Wastes. EPA-600/4-79-020, U.S. EPA-EMSL, Cincinnati, Ohio.

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# ATTACHMENT A

# GENERAL SAMPLE CONTAINER AND PRESERVATION REQUIREMENTS

Sample T	ype and Concentra	tion	Container <sup>(1)</sup>	Sample Size	Preservation <sup>(2)</sup>	Holding Time <sup>(2)</sup>
WATER	<del></del>			<u> </u>		
Organics (GC&GC/MS)	VOC	Low	Borosilicate glass	2 x 40 mL	Cool to 4°C HCl to ≤ 2	14 days <sup>(9)</sup>
	Extractables SVOCs and pesticide/PCBs)	(Low	Amber glass	2x2 L or 4x1 L	Cool to 4°C	7 days to extraction; 40 days after extraction
	Extractables SVOCs and pesticide/PCBs)	(Medium	Amber glass	2x2 L or 4x1 L	None	7 days to extraction; 40 days after extraction
Inorganics	Metals	Low	High-density polyethylene	1L	HNO <sub>3</sub> to pH ≤2	6 months (Hg-28 days
		Medium	Wide-mouth glass	16 oz.	None	6 months
	Cyanide	Low	High-density polyethylene	1 L	NaOH to pH>12	14 days
	Cyanide	Medium	Wide-mouth glass	16 oz.	None	14 days
Organic/ Inorganic	High Hazard		Wide-mouth glass	8 oz.	None	14 days
SOIL	<u> </u>		•			•
Organics (GC&GC/MS)	VOC		EnCore Sampler	(3) 5 g Samplers	Cool to 4°C	48 hours to lab preservation
	Extractables SVOCs and pesticides/PCBs)	(Low	Wide-mouth glass	8 oz.	Cool to 4°C	14 days to extraction; 40 days after extraction
	Extractables SVOCs and pesticides/PCBs)	(Medium	Wide-mouth glass	8 oz.	Cool to 4°C	14 days to extraction; 40 days after extraction
Inorganics	Low/Medium		Wide-mouth glass	8 oz.	Cool to 4°C	6 months (Hg - 28 days) Cyanide (14 days)
Organic/Inorga nic	High Hazard		Wide-mouth glass	8 oz.	None	NA
Dioxin/Furan	All		Wide-mouth glass	4 oz.	None	35 days until extraction; 40 days after extraction
TCLP	All		Wide-mouth glass	8 oz.	None	7 days until preparation; analysis as per fraction
AIR						
Volatile Organics	Low/Medium	·	Charcoal tube 7 cm long, 6 mm OD, 4 mm ID	100 L air	Cool to 4°C	5 days recommended

All glass containers should have Teflon cap liners or septa. See Attachment E. Preservation and maximum holding time allowances per 40 CFR 136.

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# **ATTACHMENT B**

# ADDITIONAL REQUIRED CONTAINERS, PRESERVATION TECHNIQUES, AND HOLDING TIMES

Parameter Number/Name	Container <sup>(1)</sup>	Preservation <sup>(2)(3)</sup>	Maximum Holding Time <sup>(4)</sup>			
INORGANIC TESTS:						
Acidity	P, G	Cool, 4°C	14 days			
Alkalinity	P, G	Cool, 4°C	14 days			
Ammonia - Nitrogen	P, G	Cool, 4°C; H₂SO₄ to pH 2	28 days			
Biochemical Oxygen Demand (BOD)	P, G	Cool, 4°C	48 hours			
Bromide	P, G	None required	28 days			
Chemical Oxygen Demand (COD)	P, G	Cool, 4°C; H <sub>2</sub> SO <sub>4</sub> to pH 2	28 days			
Chloride	P, G	None required	28 days			
Chlorine, Total Residual	P, G	None required	Analyze immediately			
Color	P, G	Cool, 4°C	48 hours			
Cyanide, Total and Amenable to Chlorination	P, G	Cool, 4°C; NaOH to pH 12; 0.6 g ascorbic acid <sup>(5)</sup>	14 days <sup>(6)</sup>			
Fluoride	Р	None required	28 days			
Hardness	P, G	HNO <sub>3</sub> to pH 2; H <sub>2</sub> SO <sub>4</sub> to pH 2	6 months			
Total Kjeldahl and Organic Nitrogen	P, G	Cool, 4°C; H <sub>2</sub> SO <sub>4</sub> to pH 2	28 days			
Nitrate - Nitrogen	P, G	None required	48 hours			
Nitrate-Nitrite - Nitrogen	P, G	Cool, 4°C; H₂SO₄ to pH 2	28 days			
Nitrite - Nitrogen	P, G	Cool, 4°C	48 hours			
Oil & Grease	G	Cool, 4°C; H <sub>2</sub> SO <sub>4</sub> to pH 2	28 days			
Total Organic Carbon (TOC)	P, G	Cool, 4°C; HCl or H <sub>2</sub> SO <sub>4</sub> to pH 2	28 days			
Orthophosphate	P, G	Filter immediately; Cool, 4°C	48 hours			
Oxygen, Dissolved-Probe	G Bottle & top	None required	Analyze immediately			
Oxygen, Dissolved-Winkler	G Bottle & top	Fix on site and store in dark	8 hours			
Phenois	G	Cool, 4°C; H₂SO₄ to pH 2	28 days			
Phosphorus, Total	P, G	Cool, 4°C; H <sub>2</sub> SO <sub>4</sub> to pH 2	28 days			
Residue, Total	P, G	Cool, 4°C	7 days			
Residue, Filterable (TDS)	P, G	Cool, 4°C	7 days			
Residue, Nonfilterable (TSS)	P, G	Cool, 4°C	7 days			
Residue, Settleable	P, G	Cool, 4°C	48 hours			
Residue, Volatile (Ash Content)	P, G	Cool, 4°C	7 days			
Silica	P	Cool, 4°C	28 days			
Specific Conductance	P, G	Cool, 4°C	28 days			
Sulfate	P, G	Cool, 4°C	28 days			

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# ATTACHMENT B ADDITIONAL REQUIRED CONTAINERS, PRESERVATION TECHNIQUES, AND HOLDING TIMES PAGE TWO

Parameter Number/Name	Container <sup>(1)</sup>	Preservation <sup>(2)(3)</sup>	Maximum Holding Time <sup>(4)</sup>
INORGANIC TESTS (Cont'd):			
Sulfide	P, G	Cool, 4°C; add zinc acetate plus sodium hydroxide to pH 9	7 days
Sulfite	P, G	None required	Analyze immediately
Turbidity	P, G	Cool, 4°C	48 hours
METALS:(7)			
Chromium VI (Hexachrome)	P, G	Cool, 4°C	24 hours
Mercury (Hg)	P, G	HNO₃ to pH 2	28 days
Metals, except Chromium VI and Mercury	P, G	HNO <sub>3</sub> to pH 2	6 months
ORGANIC TESTS:(8)			
Purgeable Halocarbons	G, Teflon-lined septum	Cool, 4°C; 0.008% Na <sub>2</sub> S <sub>2</sub> O <sub>3</sub> <sup>(5)</sup>	14 days
Purgeable Aromatic Hydrocarbons	G, Teflon-lined septum	Cool, 4°C; 0.008% Na <sub>2</sub> S <sub>2</sub> O <sub>3</sub> <sup>(5)</sup> HCl to pH 2 <sup>(9)</sup>	14 days
Acrolein and Acrylonitrile	G, Teflon-lined septum	Cool, 4°C; 0.008% Na <sub>2</sub> S <sub>2</sub> O <sub>3</sub> <sup>(5)</sup> adjust pH to 4-5 <sup>(10)</sup>	14 days
Phenois <sup>(11)</sup>	G, Teflon-lined cap	Cool, 4°C; 0.008% Na <sub>2</sub> S <sub>2</sub> O <sub>3</sub> <sup>(5)</sup>	7 days until extraction 40 days after extraction
Benzidines <sup>(11), (12)</sup>	G, Teflon-lined cap	Cool, 4°C; 0.008% Na <sub>2</sub> S <sub>2</sub> O <sub>3</sub> <sup>(5)</sup>	7 days until extraction <sup>(13)</sup>
Phthalate esters <sup>(11)</sup>	G, Teflon-lined cap	Cool, 4°C	7 days until extraction 40 days after extraction
Nitrosamines <sup>(11), (14)</sup>	G, Teflon-lined cap	Cool, 4°C; store in dark; 0.008% Na <sub>2</sub> S <sub>2</sub> O <sub>3</sub> <sup>(5)</sup>	7 days until extraction 40 days after extraction
PCBs <sup>(11)</sup>	G, Teflon-lined cap	Cool, 4°C	7 days until extraction 40 days after extraction
Nitroaromatics & Isophorone <sup>(11)</sup>	G, Teflon-lined cap	Cool, 4°C; 0.008% Na <sub>2</sub> S <sub>2</sub> O <sub>3</sub> <sup>(5)</sup> ; store in dark	7 days until extraction 40 days after extraction
Polynuclear Aromatic Hydrocarbons (PAHs) <sup>(11),(14)</sup>	G, Teflon-lined cap	Cool, 4°C; 0.008% Na <sub>2</sub> S <sub>2</sub> O <sub>3</sub> <sup>(5)</sup> ; store in dark	7 days until extraction 40 days after extraction
Haloethers <sup>(11)</sup>	G, Teflon-lined cap	Cool, 4°C; 0.008% Na <sub>2</sub> S <sub>2</sub> O <sub>3</sub> <sup>(5)</sup>	7 days until extraction 40 days after extraction
Dioxin/Furan (TCDD/TCDF) <sup>(11)</sup>	G, Teflon-lined cap	Cool, 4°C; 0.008% Na <sub>2</sub> S <sub>2</sub> O <sub>3</sub> <sup>(5)</sup>	7 days until extraction 40 days after extraction

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# ATTACHMENT B ADDITIONAL REQUIRED CONTAINERS, PRESERVATION TECHNIQUES, AND HOLDING TIMES PAGE THREE

(1) Polyethylene (P): generally 500 ml or Glass (G): generally 1L.

(2) Sample preservation should be performed immediately upon sample collection. For composite chemical samples each aliquot should be preserved at the time of collection. When use of an automated sampler makes it impossible to preserve each aliquot, then chemical samples may be preserved by maintaining at 4°C until compositing and sample splitting is completed.

(3) When any sample is to be shipped by common carrier or sent through the United States Mail, it must comply with the Department of Transportation Hazardous Materials Regulations (49 CFR Part 172).

(4) Samples should be analyzed as soon as possible after collection. The times listed are the maximum times that samples may be held before analysis and still be considered valid. Samples may be held for longer periods only if the permittee, or monitoring laboratory, has data on file to show that the specific types of samples under study are stable for the longer periods and has received a variance from the Regional Administrator.

(5) Should only be used in the presence of residual chlorine.

(6) Maximum holding time is 24 hours when sulfide is present. Optionally, all samples may be tested with lead acetate paper before pH adjustments are made to determine if sulfide is present. If sulfide is present, it can be removed by the addition of cadmium nitrate powder until a negative spot test is obtained. The sample is filtered and then NaOH is added to pH 12.

(7) Samples should be filtered immediately on site before adding preservative for dissolved metals.

(8) Guidance applies to samples to be analyzed by GC, LC, or GC/MS for specific compounds.

(9) Sample receiving no pH adjustment must be analyzed within 7 days of sampling.

(10) The pH adjustment is not required if acrolein will not be measured. Samples for acrolein receiving no pH adjustment must be analyzed within 3 days of sampling.

- (11) When the extractable analytes of concern fall within a single chemical category, the specified preservative and maximum holding times should be observed for optimum safeguard of sample integrity. When the analytes of concern fall within two or more chemical categories, the sample may be preserved by cooling to 4°C, reducing residual chlorine with 0.008% sodium thiosulfate, storing in the dark, and adjusting the pH to 6-9; samples preserved in this manner may be held for 7 days before extraction and for 40 days after extraction. Exceptions to this optional preservation and holding time procedure are noted in footnote 5 (re: the requirement for thiosulfate reduction of residual chlorine) and footnotes 12, 13 (re: the analysis of benzidine).
- (12) If 1,2-diphenylthydrazine is likely to be present, adjust the pH of the sample to 4.0±0.2 to prevent rearrangement to benzidine.
- (13) Extracts may be stored up to 7 days before analysis if storage is conducted under an inert (oxidant-free) atmosphere.
- (14) For the analysis of diphenylnitrosamine, add 0.008% Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub> and adjust pH to 7-10 with NaOH within 24 hours of sampling.
- (15) The pH adjustment may be performed upon receipt at the laboratory and may be omitted if the samples are extracted within 72 hours of collection. For the analysis of aldrin, add 0.008% Na₂S₂O₃.



**TETRA TECH NUS, INC.** 

# STANDARD OPERATING PROCEDURES

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Applicability

Tetra Tech NUS, Inc.

Prepared

Earth Sciences Department

Subject

FIELD DOCUMENTATION

Approved

Tom Johnston



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ATTAG	HMENTS
A B C	TYPICAL SITE LOGBOOK ENTRY

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### 1.0 PURPOSE

The purpose of this Standard Operating Procedure (SOP) is to identify and designate the field data record forms, logs, and reports generally initiated and maintained for documenting Tetra Tech NUS, Inc. (TtNUS) field activities.

## 2.0 SCOPE

Documents presented within this SOP (or equivalents) shall be used for all TtNUS field activities, as applicable. Other or additional documents may be required by specific client contracts or project planning documents.

#### 3.0 GLOSSARY

None.

# 4.0 RESPONSIBILITIES AND PERSONNEL QUALIFICATIONS

<u>Project Manager (PM)</u> - The PM is responsible for obtaining hardbound controlled-distribution logbooks (from the appropriate source), as needed. In addition, the Project Manager is responsible for placing all field documentation used in site activities (i.e., records, field reports, sample data sheets, field notebooks, and the site logbook) in the project's central file upon the completion of field work.

<u>Field Operations Leader (FOL)</u> - The FOL is responsible for ensuring that the site logbook, notebooks, and all appropriate and current forms and field reports included in this SOP (and any additional forms required by the contract) are correctly used, accurately filled out, and completed in the required time frame.

General personnel qualifications for field documentation activities include the following:

- Occupational Safety and Health Administration (OSHA) 40-hour and applicable refresher training.
- Capability of performing field work under the expected physical and environmental (i.e., weather)
  conditions.
- Familiarity with appropriate procedures for documentation, handling, packaging, and shipping.

# 5.0 PROCEDURES

## 5.1 SITE LOGBOOK

# 5.1.1 General

The site logbook is a hard-bound, paginated, controlled-distribution record book in which all major on-site activities are documented. At a minimum, record or reference the following activities/events (daily) in the site logbook:

- All field personnel present
- Arrival/departure times and names of site visitors
- Times and dates of health and safety training
- Arrival/departure times of equipment
- Times and dates of equipment calibration

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- Start and/or completion of borehole, trench, monitoring well installation activities, etc.
- Daily on-site activities
- Sample pickup information
- Health and safety issues (level of protection, personal protective equipment [PPE], etc.)
- Weather conditions

Maintain a site logbook for each project and initiate it at the start of the first on-site activity (e.g., site visit or initial reconnaissance survey). Make entries every day that on-site activities take place involving TtNUS or subcontractor personnel. Upon completion of the fieldwork, provide the site logbook to the PM or designee for inclusion in the project's central file.

Record the following information on the cover of each site logbook:

- Project name
- TtNUS project number
- Sequential book number
- Start date
- End date

Information recorded daily in the site logbook need not be duplicated in other field notebooks (see Section 5.2) but must summarize the contents of these other notebooks and refer to specific page locations in these notebooks for detailed information (where applicable). An example of a typical site logbook entry is shown in Attachment A.

If measurements are made at any location, either record the measurements and equipment used in the site logbook or reference the field notebook in which the measurements are recorded (see Attachment A).

Make all logbook, notebook, and log sheet entries in indelible ink (black pen is preferred). No erasures are permitted. If an incorrect entry is made, cross out the entry with a single strike mark, initial, and date it. At the completion of entries by any individual, the logbook pages used must be signed and dated by the person making the entries. The site logbook must also be signed by the FOL at the end of each day.

### 5.1.2 Photographs

Sequentially number movies, slides, or photographs taken of a site or any monitoring location to correspond to logbook/notebook entries. Enter the name of the photographer, date, time, site location, site description, and weather conditions in the logbook/notebook as the photographs are taken. A series entry may be used for rapid-sequence photographs. The photographer is not required to record the aperture settings and shutter speeds for photographs taken within the normal automatic exposure range. However, special lenses, films, filters, and other image-enhancement techniques must be noted in the logbook/notebook. If possible, such techniques shall be avoided because they can adversely affect the accuracy of photographs. Chain-of-custody procedures depend on the subject matter, type of camera (digital or film), and the processing it requires. Follow chain-of-custody procedures for film used for aerial photography, confidential information, or criminal investigation. After processed, consecutively number the slides of photographic prints and label them according to the logbook/notebook descriptions. Docket the site photographs and associated negatives and/or digitally saved images to compact disks into the project's central file.

# 5.2 FIELD NOTEBOOKS

Key field team personnel may maintain a separate dedicated field notebook to document the pertinent field activities conducted directly under their supervision. For example, on large projects with multiple investigative sites and varying operating conditions, the Health and Safety Officer may elect to maintain a

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separate field notebook. Where several drill rigs are in operation simultaneously, each site geologist assigned to oversee a rig must maintain a field notebook.

# 5.3 FIELD FORMS

All TtNUS field forms (see list in Section 6.0 of this SOP) can be found on the company's intranet site (<a href="http://intranet.ttnus.com">http://intranet.ttnus.com</a>) under Field Log Sheets. Forms may be altered or revised for project-specific needs, subject to client approval. Care must be taken to ensure that all essential information can be documented. Guidelines for completing these forms can be found in the related sampling SOPs.

# 5.3.1 Sample Collection, Labeling, Shipment, Request for Analysis, and Field Test Results

# 5.3.1.1 Sample Log Sheet

Sample log sheets are used to record specified types of data while sampling. The data recorded on these sheets are useful in describing the sample as well as pointing out any problems, difficulties, or irregularities encountered during sampling. Complete a sample log sheet for each sample obtained, including field quality control (QC) samples.

### 5.3.1.2 Sample Label

A typical sample label is illustrated in Attachment B. Complete the required information on the adhesive labels and apply them to every sample container. Obtain sample labels from the appropriate program/project source, request that they be electronically generated in house, or request them the laboratory subcontractor.

# 5.3.1.3 Chain-of-Custody Record

The chain-of-custody record is a multi-part form that is initiated as samples are acquired and accompanies a sample (or group of samples) as they are transferred from person to person. This form must be used as follows for any samples collected for chemical or geotechnical analysis whether the analyses are performed on site or off site:

- Retain one carbonless copy of the completed chain-of custody form in the field.
- Send one copy is sent to the PM (or designee)
- Send the original to the laboratory with the associated samples. Place the original (top, signed copy) of the chain-of custody form inside a large Ziploc®-type bag taped inside the lid of the shipping cooler. If multiple coolers are sent but are included on one chain-of custody form, send the form with the cooler containing vials for volatile organic compound (VOC) analysis or the cooler with the air bill attached. Indicate on the air bill how many coolers are included with that shipment.

An example of a chain-of-custody form is provided as Attachment C. After the samples are received at the laboratory, the sample cooler and contents are checked and any problems are noted on the enclosed chain-of custody form (any discrepancies between the sample labels and chain-of custody form and any other problems that are noted are resolved through communication between the laboratory point-of-contact and the TtNUS PM). The chain-of custody form is signed and copied. The laboratory will retain the copy, and the original becomes part of the samples' corresponding analytical data package.

## 5.3.1.4 Chain-of-Custody Seal

Attachment D is an example of a custody seal. The custody seal is an adhesive-backed label that is part of a chain-of-custody process and is used to prevent tampering with samples after they have been collected in the field and sealed in coolers for transport to the laboratory. Sign and date custody seals

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and affix them across the lid and body of each cooler (front and back) containing environmental samples (see SOP SA-6.1). Obtain custody seals from the laboratory (if available) or purchase them from a supplier.

## 5.3.1.5 Geochemical Parameters Log Sheets

Complete Field Analytical Log Sheets to record geochemical and/or natural attenuation field test results.

# 5.3.2 Hydrogeological and Geotechnical Forms

# 5.3.2.1 Groundwater Level Measurement Sheet

Complete a Groundwater Level Measurement Sheet for each round of water level measurements made at a site.

# 5.3.2.2 Data Sheet for Pumping Test

During the performance of a pumping test (or an in-situ hydraulic conductivity test), a large amount of data must be recorded, often within a short time period. Use a Pumping Test Data Sheet to facilitate this task-by standardizing the data collection format for the pumping-well-and observation wells, and allowing the time interval for collection to be established in advance.

### 5.3.2.3 Packer Test-Report-Form

Complete a Packer Test Report Form for each-well at which a packer test is conducted.

## 5.3.2.4 Boring Log

Complete a Summary Log of Boring, or Boring Log for each soil boring performed to document the materials encountered, operation and driving of casing, and locations/depths of samples collected. In addition, if volatile organics are monitored on cores, samples, cuttings from the borehole, or breathing zone, (using a photoionization detector [PID] or flame ionization detector [FID]), enter these readings on the boring log at the appropriate depth. When they become available, enter the laboratory sample number, concentrations of key contaminants, or other pertinent information in the "Remarks" column. This feature allows direct comparison of contaminant concentrations with soil characteristics.

## 5.3.2.5 Monitoring Well Construction Details Form

Complete a Monitoring Well Construction Details Form for every monitoring well, piezometer, or temporary well point installed. This form contains specific information on length and type of well riser pipe and screen, backfill, filter pack, annular seal and grout characteristics, and surface seal characteristics. This information is important in evaluating the performance of the monitoring well, particularly in areas where water levels show temporal variation or where there are multiple (immiscible) phases of contaminants. Depending on the type of monitoring well (in overburden or bedrock, stick-up or flush mount), different forms are used.

# 5.3.2.6 <u>Test Pit Log</u>

When a test pit or trench is constructed for investigative or sampling purposes, a Test Pit Log must be filled out by the responsible field geologist or sampling technician.

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# 5.3.2.7 <u>Miscellaneous Monitoring Well Forms</u>

Miscellaneous monitoring well forms that may be required on a project-specific basis include the Monitoring Well Materials Certificate of Conformance and Monitoring Well Development Record. Use a Monitoring Well Materials Certificate of Conformance to document all materials utilized during each monitoring well installation. Use a Monitoring Well Development Record to document all well development activities.

# 5.3.2.8 <u>Miscellaneous Field Forms</u> – Quality Assurance and Checklists

Miscellaneous field forms/checklists forms that may be required on a project-specific basis include the following:

- Container Sample and Inspection Sheet use this form when a container (drum, tank, etc.) is sampled and/or inspected.
- QA Sample Log Sheet use this form when a QA sample such as an equipment rinsate blank, source blank, etc. is collected.
- Field Task Modification Request (FTMR) use this form to document deviations from the project planning documents. The FOL is responsible for initiating the FTMRs. Maintain copies of all FTMRs with the on-site planning documents, and place originals in the final evidence file.
- Field Project Daily Activities Checklist and Field Project Pre-Mobilization Checklist used these during both the planning and field effort to ensure that all necessary tasks are planned for and completed. These two forms are not requirements but are useful tools for most field work.

# 5.3.3 Equipment Calibration and Maintenance Form

The calibration or standardization of monitoring, measuring, or test equipment is necessary to ensure the proper operation and response of the equipment, to document the accuracy, precision, or sensitivity of the measurements, and determine if correction should be applied to the readings. Some items of equipment require frequent calibration, others infrequent. Some are calibrated by the manufacturer, others by the user.

Each instrument requiring calibration has its own Equipment Calibration Log, which documents that the manufacturer's instructions were followed for calibration of the equipment, including frequency and type of standard or calibration device. Maintain an Equipment Calibration Log for each electronic measuring device used in the field; make entries for each day the equipment is used or in accordance with manufacturer recommendations.

### 5.4 FIELD REPORTS

The primary means of recording on-site activities is the site logbook. Other field notebooks may also be maintained. These logbooks and notebooks (and supporting forms) contain detailed information required for data interpretation or documentation but are not easily used for tracking and reporting of progress. Furthermore, the field logbook/notebooks remain on site for extended periods of time and are thus not accessible for timely review by project management. Other reports useful for tracking and reporting the progress of field activities are described below.

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# 5.4.1 Daily Activities Report

To provide timely oversight of on-site contractors, complete and submit Daily Activities Reports (DARs) as described below.

# 5.4.1.1 Description

The DAR documents the activities and progress for each day's field work. Complete this report on a daily basis whenever there are drilling, test pitting, well construction, or other related activities occurring that involve subcontractor personnel. These sheets summarize the work performed and form the basis of payment to subcontractors. The DAR form can be found on the TtNUS intranet site.

### 5.4.1.2 Responsibilities

It is the responsibility of the rig geologist to complete the DAR and obtain the driller's signature acknowledging that the times and quantities of material entered are correct.

### 5.4.1.3 Submittal and Approval

At the end of the shift, the rig geologist must submit the DAR to the FOL for review and filing. The Daily Activities Report is not a formal report and thus requires no further approval. The DARs are retained by the FOL for use in preparing the site logbook and in preparing weekly status reports for submission to the PM.

## 5.4.2 Weekly Status Reports

To facilitate timely review by project management, photocopies of logbook/notebook entries may be made for internal use.

In addition to those described herein, other summary reports may also be contractually required.

All TtNUS field forms can be found on the company's intranet site at <a href="http://intranet.ttnus.com">http://intranet.ttnus.com</a> under Field Log Sheets.

# 6.0 LISTING OF FIELD FORMS ON THE TUNUS INTRANET SITE

- Boring Log
- Container Sample and Inspection Sheet
- Daily Activities Checklist
- Daily Activities Record
- Equipment Calibration Log
- Field Task Modification Request
- Field Analytical Log sheet Geochemical Parameters
- Groundwater Level Measurement Sheet
- Groundwater Sample Log Sheet
- Hydraulic Conductivity Test Data Sheet
- Low Flow Purge Data Sheet
- Bedrock Monitoring Well Construction (Stick Up)
- Bedrock Monitoring Well Construction Flush Mount
- Bedrock Monitoring Well Construction Open Hole
- Confining Layer Monitoring Well Construction
- Monitoring Well Development Record

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- Monitoring Well Materials Certificate of Conformance
- Overburden Monitoring Well Construction Flush Mount
- Overburden Monitoring Well Construction Stick Up
- Packer Test Report Form
- Pumping Test Data Sheet
- QA Sample Log Sheet
- Soil/Sediment Sample Log Sheet
- Surface Water Sample Log Sheet
- Test Pit Log
- Field Project Pre-Mobilization Checklist

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# ATTACHMENT A TYPICAL SITE LOGBOOK ENTRY

START 1	TIME: DATE:
SITE LE	
PERSON	NEL: TtNUS DRILLER SITE VISITORS
WEATHE	ER: Clear, 68°F, 2-5 mph wind from SE
ACTIVITI	ES:
1.	Steam jenney and fire hoses were set up.
2.	Drilling activities at well resumes. Rig geologist was Se Geologist's Notebook, No. 1, page 29-30, for details of drilling activity. Sample No. 123-2' S4 collected; see sample logbook, page 42. Drilling activities completed at 11:50 and 4-inch stainless steel well installed. See Geologist's Notebook, No. 1, page 31, and we construction details for well
3.	Drilling rig No. 2 steam-cleaned at decontamination pit. Then set up at location well
4.	Well drilled. Rig geologist was See Geologist's Notebook No. 2, page for details of drilling activities. Sample numbers 123-22-S1, 123-22-S2 and 123-22-S3 collected; see sample logbook, pages 43, 44, and 45.
5.	Well was developed. Seven 55-gallon drums were filled in the flushing stage. The well was then pumped using the pitcher pump for 1 hour. At the end of the hour, water pumped from well was "sand free."
6.	EPA remedial project manger arrives on site at 14:25 hours.
7.	Large dump truck arrives at 14:45 and is steam-cleaned. Backhoe and dump truck set u over test pit
8.	Test pit dug with cuttings placed in dump truck. Rig geologist wa See Geologist's Notebook, No. 1, page 32, for details of test pactivities. Test pit subsequently filled. No samples taken for chemical analysis. Due to shallow groundwater table, filling in of test pit resulted in a very soft and wet area. mound was developed and the area roped off.
9.	Express carrier picked up samples (see Sample Logbook, pages 42 through 45) a 17:50 hours. Site activities terminated at 18:22 hours. All personnel off site, gate locked.
	Field Operations Leader

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## ATTACHMENT B SAMPLE LABEL

TŁ.	Tetra Tech 661 Anders Pittsburgh, (412)921-70	en Drive 15220	Project: Site: Location:	
Sample N	lo:			Matrix:
Date:		Time:	Preserve	e:
Analysis	•	ï		1
Sampled	by:		Laborato	ory:

Number

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## STANDARD OPERATING PROCEDURES

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Applicability

Tetra Tech NUS, Inc.

Prepared

Earth Sciences Department

Subject DECONTAMINATION OF FIELD EQUIPMENT

Approved

Tom Johnston



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#### 1.0 PURPOSE

Decontamination is the process of removing and/or neutralizing site contaminants that have contacted and/or accumulated on equipment. The purpose of this Standard Operating Procedure (SOP) is to protect site personnel, the general public, and the environment while preserving or maintaining sample integrity. It is further intended through this procedure to describe the steps necessary for proper decontamination of drilling equipment, earth-moving equipment, chemical sampling equipment and field operation and analytical equipment.

#### 2.0 SCOPE AND APPLICABILITY

This procedure applies to all equipment used to provide access to/acquire environmental samples that may have become contaminated through direct contact with contaminated media including air, water, and soil. This equipment includes drilling and heavy equipment and chemical sampling and field analytical equipment. Where technologically and economically feasible, single-use sealed disposable equipment will be employed to minimize the potential for cross-contamination. This SOP also provides general reference information on the control of contaminated materials.

Decontamination methods and equipment requirements may differ from one project to another. General equipment items are specified in Section 6.0, but project-specific equipment must be obtained to address the project-specific decontamination procedures presented in Section 7.0 and applicable subsections.

#### 3.0 GLOSSARY

Alconox/Liquinox - A brand of phosphate-free laboratory-grade detergent.

<u>Decontamination Solution</u> - A solution selected/identified in the Health and Safety Plan or Project-Specific Quality Assurance Plan. The solution is selected and employed as directed by the project chemist/health and safety professional.

<u>Deionized Water (DI)</u> - Tap water that has been treated by passing through a standard deionizing resin column. This water may also pass through additional filtering media to attain various levels of analyte-free status. The DI water should meet College of American Pathologists (CAP) and National Committee for Clinical Laboratory Standards (NCCLS) specifications for reagent-grade Type I water.

<u>Potable Water</u> - Tap water from any municipal water treatment system. Use of an untreated potable water supply is not an acceptable substitute for tap water.

<u>Pressure Washing</u> - Process employing a high-pressure pump and nozzle configuration to create a high-pressure spray of potable water. High-pressure spray is employed to remove solids from equipment.

<u>Solvent</u> – A liquid in which solid chemicals or other liquids are dissolved. The solvent of choice is pesticide-grade isopropanol. Use of other solvents (methanol, acetone, or hexane) may be required for particular projects or for a particular purpose (e.g., removal of concentrated waste) and must be justified in the project planning documents. For example, it may be necessary to use hexane when analyzing for trace levels of pesticides, PCBs, or fuels. In addition, because many of these solvents are not miscible in water, the equipment should be air dried prior to use. Solvents should not be used on PVC equipment or well construction materials.

<u>Steam Pressure Washing</u> - A cleaning method employing a high-pressure spray of heated potable water to remove various organic/inorganic chemicals from equipment.

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#### 4.0 RESPONSIBILITIES AND PERSONNEL QUALIFICATIONS

<u>Project Manager</u> - Responsible for ensuring that all field activities are conducted in accordance with approved project plan(s) requirements.

<u>Decontamination Personnel</u> - Individuals assigned the task of decontamination. It is the responsibility of these individuals to understand the use and application of the decontamination process and solutions as well as the monitoring of that process to ensure that it is working properly. This is accomplished through visual evaluation, monitoring instrument scanning of decontaminated items, and/or through the collection of rinsate blanks to verify contaminant removal.

<u>Field Operations Leader (FOL)</u> - Responsible for the implementation of project-specific planning documents. This includes on-site verification that all field activities are performed in compliance with approved SOPs or as otherwise dictated by the approved project plan(s). The FOL is also responsible for the completion and accuracy of all field documentation.

<u>Site Safety Officer (SSO)</u> - Exercises shared responsibility with the FOL concerning decontamination effectiveness. All equipment arriving on site (as part of the equipment inspection), leaving the site, and moving between locations is required to go through a decontamination evaluation. This is accomplished through visual examination and/or instrument screening to determine the effectiveness of the decontamination process. Improper or incomplete decontamination is sufficient to restrict equipment from entering the site, exiting the site, or moving to a new location on the site until the objectives are successfully completed.

General personnel qualifications for decontamination activities include the following:

- Occupational Safety and Health Administration (OSHA) 40-hour and applicable refresher training.
- Capability of performing field work under the expected physical and environmental (i.e., weather)
  conditions.
- Familiarity with appropriate decontamination procedures.

#### 5.0 HEALTH AND SAFETY

In addition to the health and safety issues and reminders specified in subsections of this SOP, the following considerations and requirements must be observed as SOPs for field equipment decontamination activities:

- If any solvents or hazardous chemicals (e.g., isopropyl alcohol) are to be used in equipment
  decontamination activities, the FOL must first obtain the manufacturer's/supplier's Material Safety
  Data Sheet (MSDS) and assure that it is reviewed by all users (prior to its use), added to the site
  Hazardous Chemical Inventory, and maintained on site as part of the project Hazard Communication
  Program.
- Review and observe specific health and safety requirements (e.g., personal protective equipment [PPE]) specified in the project-specific health and safety plan for this activity.

#### 6.0 EQUIPMENT LIST

Wood for decontamination pad construction, when applicable (see Section 7.1).

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- Tools for constructing decontamination pad frame, when applicable (see Section 7.1).
- Visqueen sheeting or comparable material to cover decontamination pad frame, when applicable (see Section 7.1).
- Wash/drying racks for auger flights and drill/drive rods, when applicable (see Section 7.2).
- PPE as specified in the project health and safety plan.
- Soap and water for washing and rinsing.
- Deionized water for final rinsing.
- Solvents (e.g., pesticide-grade isopropanol) for rinsing (see applicable portions of Section 7.2).
- Tubs, buckets, etc. for containerizing rinse water (see applicable portions of Section 7.2).
- Sample bottles for collecting rinsate blanks (see Section 7.2).
- Calibrated photoionization detector (PID) or flame ionization detector (FID) to monitor decontaminated equipment for organic vapors generated through the existence of residual contamination or the presence of decontamination solvent remaining after the piece was rinsed.
- Aluminum foil or clear clean plastic bag for covering cleaned equipment (see applicable portions of Section 7.2).
- Paper towels or cloths for wiping.
- Brushes, scrapers, or other hand tools useful for removing solid materials from equipment.
- Clear plastic wrap for covering or wrapping large decontaminated equipment items (see Section 7.2.2).
- Drum-moving equipment for moving filled waste drums (optional) (see Section 7.3).
- Drum labels for waste drums (see Attachment A).

#### 7.0 PROCEDURES

The process of decontamination is accomplished through the removal of contaminants, neutralization of contaminants, or isolation of contaminants. To accomplish this activity, preparation is required including site preparation, equipment selection, and evaluation of the decontamination requirements and processes. Site contaminant types, concentrations, and media types are primary drivers in the selection of the types of decontamination and where it will be conducted. For purposes of this SOP, discussion is limited to decontamination procedures for general environmental investigations.

Decontamination processes will be performed at the location(s) specified in project-specific planning documents. Typical decontamination locations include the following:

- Temporary decontamination pads/facilities
- Sample locations
- Centralized decontamination pad/facilities

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Combination of some or all of the above

The following discussion includes general considerations for the decontamination process. Specific construction and implementation procedures will be as specified in the project-specific planning documents and/or may be as dictated by site-specific conditions as long as the intent of the requirements in the planning documents is met. This intent is to contain any residual fluids and solids generated through the decontamination process.

#### 7.1 <u>Decontamination Pad Design/Construction Considerations</u>

#### 7.1.1 Temporary Decontamination Pads

Temporary decontamination pads may be constructed at satellite locations within the site area in support of temporary work areas. These structures are generally constructed to support the decontamination of heavy equipment such as drill rigs and earth-moving equipment but can be employed for smaller articles.

The purpose of the decontamination pad is to contain wash waters and potentially contaminated soil generated during decontamination procedures. Therefore, construction of these pads should take into account the following considerations:

- Site location The decontamination site selected should be far enough from the work site to maximize decontamination effectiveness while minimizing travel distance. The location of the decontamination site shall be selected to provide, in the judgment of the FOL or FOL designee, compliance with as many of the following characteristics as practicable:
  - Well removed from pedestrian/vehicle thoroughfares.
  - Avoidance of areas where control/custody cannot be maintained.
  - Avoidance of areas where potential releases of contaminated media or decontamination fluids may be compounded through access to storm water transport systems, streams, or other potentially sensitive areas.
  - Avoidance of potentially contaminated areas.
  - Avoidance of areas too close to the ongoing operation, where cross-contamination may occur.

The selected decontamination site should include the following, where possible:

- Areas where potable water and electricity are provided.

#### Safety Reminder

When utilizing electrical power sources, either hard-wired or portable-generated sources, ensure that:

- All power is routed through a Ground Fault Circuit Interrupter (GFCI).
- All power cords are in good condition (no physical damage), rated for the intended energy load, and designated for outdoor use.

In situations where accomplishing these elements is not possible, it will be necessary to implement a site electrical grounding program.

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- Areas where support activities such as removing decontamination waters soil and sediment are possible without entering an active exclusion zone.
- Areas that offer sufficient size to carry out the specific decontamination sequence.
- Decontamination pad (decon pad) The decon pad shall be constructed to meet the following characteristics:
  - Size The size of the pad should be sufficient to accept the equipment to be decontaminated as well as permitting free movement around the equipment by the personnel conducting the decontamination. The size should permit these movements utilizing pressure/steam washer wands and hoses and minimizing splash due to work in close quarters.
  - Slope An adequate slope will be constructed to permit the collection of water and potentially contaminated soil within a trough or sump constructed at one end. The collection point for wash waters should be of adequate distance that the decontamination workers do not have to walk through the wash waters while completing their tasks. Because the pad will be sloped, place a light coating of sand over the plastic to minimize potential slips and falls. See the text about liners below.
  - Sidewalls The sidewalls shall be at least 6 inches in height (or as high as possible if 6 inches is not achievable) to provide adequate containment for wash waters and soil. If splash represents a potential problem, splash guards should be constructed to control overspray. Sidewalls may be constructed of wood, inflatables, sand bags, etc. to permit containment. Splash guards are typically wood frames with Visqueen coverings to control overspray.
  - Liner Depending on the types of equipment and decontamination method to be used, the liner should be of sufficient thickness to provide a puncture-resistant barrier between the decontamination operation and the unprotected environment. Care should be taken to examine the surface area prior to placing the liner to remove sharp articles (sticks, stones, debris) that could puncture the liner. Liners are intended to form an impermeable barrier. The thickness may vary from a minimum recommended thickness of 10 mil to 30 mil. The desired thickness may be achieved through layering materials of lighter construction. It should be noted that various materials (rubber, polyethylene sheeting) become slippery when wet. To minimize this potential hazard associated with a sloped liner, a light coating of sand shall be applied to provide traction as necessary.
  - Wash/drying racks Auger flights, drill/drive rods, and similar equipment require racks positioned off of the ground to permit these articles to be washed, drained, and dried while secured from falling during this process.

For decontamination of direct-push technology (DPT) equipment, the pad may be as simple as a mortar tub containing buckets of soapy water for washing and an empty bucket to capture rinse waters. Decontamination may be conducted at the rear of the rig to permit rapid tool exchange.

- Maintenance Maintain the decontamination area by:
  - Periodically clearing the work area of standing water, soil, and debris, and coiling hoses to aid in eliminating slip, trip, and fall hazards. In addition, these articles will reduce potential backsplash and cross-contamination.

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- Regularly changing the decontamination fluids to ensure proper cleaning and prevent cross-contamination.
- PPE Periodically evaluate the condition of, and maintain the decontamination equipment, including regular cleaning of face shields and safety glasses. This is critical to ensuring the safety of decontamination personnel and the integrity of the decontamination process, and it will ensure that equipment is functioning properly.

#### 7.1.2 Decontamination Activities at Drill Rigs/DPT Units

During subsurface sampling activities including drilling and DPT activities, decontamination of drive rods, Macro Core Samplers, split spoons, etc. is typically conducted at an area adjacent to the operation. Decontamination is generally accomplished using a soap/water wash and rinse utilizing buckets and brushes. This area requires sufficient preparation to accomplish the decontamination objectives.

Buckets shall be placed within mortar tubs or similar secondary containment tubs to prevent splash and spills from reaching unprotected environmental media. Drying racks shall be employed as directed for temporary pads to permit parts to dry and be evaluated prior to use/reuse. Methodology regarding this activity is provided in Section 7.2.

#### 7.1.3 Decontamination Activities at Remote Sample Locations

When sampling at remote locations, sampling equipment such as trowels and pumps/tubing should be evacuated of potentially contaminated media to the extent possible. This equipment should be wrapped in plastic for transport to the temporary/centralized decontamination location for final cleaning and disposition. Flushing and cleaning of single-use equipment such as disposable trowels, tubing, and surgeon's gloves may allow disposal of this equipment after visible soil and water remnants have been removed.

#### 7.2 <u>Equipment Decontamination Procedures</u>

The following represents procedures to be employed for the decontamination of equipment that may have contacted and/or accumulated contamination through site investigation activities.

#### 7.2.1 Monitoring Well Sampling Equipment

- 7.2.1.1 <u>Groundwater sampling equipment This includes pumps inserted into monitoring wells such as bladder pumps, Whale pumps, and Redi-Flo pumps and reusable bailers, etc.</u>
- 1. Evacuate to the extent possible, any purge water within the pump/bailer.
- 2. Scrub using soap and water and/or steam clean the outside of the pump/bailer and, if applicable, the pump tubing.
- 3. Insert the pump and tubing/bailer into a clean container of soapy water. Pump/run a sufficient amount of soapy water through the pump/bailer to flush out any residual well water. After the pump is flushed, circulate soapy water through the pump to ensure that the internal components are thoroughly flushed.
- 4. Remove the pump and tubing/bailer from the container
- 5. Rinse external pump components using tap water.

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6. Insert the pump and tubing/bailer into a clean container of tap water. Pump/run a sufficient amount of tap water through the pump/bailer to evacuate all of the soapy water (until clear).

#### **CAUTION**

Do not rinse PE, PVC, and associated tubing with solvents –
Use the procedures defined in the project-specific planning documents. If they are not defined, contact the FOL for guidance. The solvent rinse described in Step 7 may be omitted if groundwater does not contain oil, grease, PAHs, PCBs, or other hard to remove organic materials.

- 7. If groundwater contains or is suspected to contain oil, grease, PAHs, PCBs, or other hard to remove organic materials, rinse the equipment to be cleaned with pesticide-grade isopropanol.
- 8. Pass deionized water through the hose to flush out the tap water and solvent residue as applicable.
- 9. Drain residual deionized water to the extent possible.
- 10. Allow components of the equipment to air dry.
- 11. For bladder pumps, disassemble the pump and wash the internal components with soap and water, then rinse with tap water, isopropanol, and deionized water and allow to dry. After the parts are dry, conduct a visual inspection and a monitoring instrument scan to ensure that potential contaminants and all decontamination solvent have been removed. Collect a rinsate blank in accordance with the project-specific planning documents to ensure that the decontamination process is functioning as intended. The typical frequency of collection for rinsate blanks is 1 per 20 field samples. In addition, wipe samples or field tests such as UV light may be used.
- 12. Wrap pump/bailer in aluminum foil or a clear clean plastic bag for storage.

#### SAFETY REMINDER

Remember when handling powered equipment to disconnect the power source and render the equipment to a zero energy state (both potential and kinetic) before opening valves, disconnecting lines, etc.

#### 7.2.1.2 Electronic Water Level Indicators/Sounders/Tapes

During water level measurements, rinsing the extracted tape and probe with deionized water and wiping the surface of the extracted tape between locations is acceptable. However, periodic full decontamination should be conducted as follows:

- 1. Wash with soap and water
- 2. Rinse with tap water
- 3. Rinse with deionized water

#### **NOTE**

In situations where oil, grease, free product, other hard to remove materials are encountered, probes and exposed tapes should be washed in hot soapy water. If probes or tapes cannot be satisfactorily decontaminated (they are still stained, discolored, etc.), they should be removed from service.

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#### 7.2.1.3 <u>Miscellaneous Equipment</u>

Miscellaneous equipment including analytical equipment (water quality testing equipment) shall be cleaned per manufacturers' instructions. This generally includes wiping the sensor housing and rinsing with tap and deionized water.

Coolers/shipping containers employed to ship samples are received from the laboratory in a variety of conditions including marginal to extremely poor. Coolers shall be evaluated prior to use for the following:

- Structural integrity Coolers missing handles or having breaks in the outer housing should be removed and not used. Notify the laboratory that the risk of shipping samples in the cooler(s) provided is too great and request a replacement unit.
- Cleanliness As per protocol, only volatile organic samples are accompanied by a trip blank. If a
  cooler's cleanliness is in question (visibly dirty/stained) or if there are noticeable odors, the cooler
  should be decontaminated prior to use as follows:
  - 1. Wash with soap and water
  - 2. Rinse with tap water
  - 3. Dry

If these measures fail to clean the cooler to an acceptable level, remove the unit from use as a shipping container and ask the cooler provider (e.g., the analytical laboratory) to provide a replacement unit.

#### 7.2.2 Downhole Drilling Equipment

This includes any portion of the drill rig that is over the borehole, including auger flights, drill stems, rods, and associated tooling that would extend over the borehole. The following procedure is to be employed prior to initiating the drilling/sampling activity, then between locations:

#### **CAUTION**

Exercise care when using scrapers to remove soil and debris from downhole drilling equipment. Inadvertent slips of scrapers have resulted in cuts, scrapes, and injured knuckles, so use scrapers carefully when removing soil from these items.

- 1. Remove loose soil using shovels, scrapers, etc.
- 2. Through a combination of scrubbing using soap and water and/or steam cleaning or pressure washing, remove visible dirt/soil from the equipment being decontaminated.

#### **CAUTION**

In Step 3, do not rinse PE, PVC, and associated tubing with solvents. The appropriate procedures should be defined within the project-specific planning documents. If they are not defined, contact the FOL for guidance. The solvent rinse described in Step 4 may be omitted if groundwater does not contain oil, grease, PAHs, PCBs, or other hard to remove organic materials.

3. Rinse the equipment with tap water, where applicable (steam cleaning and pressure washing incorporate rinsing as part of the process).

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- 4. If the equipment has directly or indirectly contacted contaminated sample media and is known or suspected of being contaminated with oil, grease, PAHs, PCBs, or other hard to remove organic materials, rinse equipment with pesticide-grade isopropanol
- 5. To the extent possible, allow components to air dry.
- If the decontaminated equipment is to be used immediately after decontamination, screen it with a
  calibrated photoionization detector (PID)/flame ionization detector (FID) to ensure that all
  contaminants and possible decontamination solvents (if they were used) have been adequately
  removed.
- 7. Wrap or cover equipment in clear plastic until it is time to be used.

#### SAFETY REMINDER

Even when equipment is disconnected from power sources, dangers such as the following may persist:

- <u>Falls</u> An auger flight standing on its end may fall and injure someone. Secure all loose articles to prevent heavy articles from falling onto people or equipment.
- <u>Burns</u> Steam cleaner water is heated to more than 212 °F and exhibits thermal energy that can cause burns. Prevent contact of skin with hot water or surfaces.

<u>High water pressure</u> - Pressure washer discharge can have 2,000 to 4,000 psi of water pressure. Water under this amount of pressure can rupture skin and other human tissues. Water at 4,000 psi exiting a 0° tip can be dangerous because of its relatively high cutting power. The exit velocity and cutting power of the water are reduced when exiting a 40° fan tip, but damage to soft tissues is still possible.

In general, follow the rules below to avoid injury, equipment damage, or incomplete decontamination:

- 1. Read the operating manual and follow the manufacturers' recommended safety practices before operating pressure washers and steam cleaners.
- Never point the pressure washer or steam cleaner at another person or use to clean your boots or other parts of your body. Water lacerations and burns may appear to be minor at first but can be life threatening. Do not attempt to hold small parts in your hand while washing them with hightemperature or high-pressure water.
- 3. Always wear PPE as specified in the HASP such as:
  - Hard hat, safety glasses, splash shield, impermeable apron or splash suit, and hearing protection. Remember that excessive noise is a hazard when operating gas-powered engines and electrically driven pressure washers. PPE will be identified in your project specific planning documents.
- 4. Inspect each device before use. An inspection checklist will be provided in the project-specific planning documents. If it is a rented device, safety measures are typically provided by the vendor. In all cases, if you are not familiar with the operation of a pressure washer/steam cleaner, do not operate it until you obtain and thoroughly review operating instructions and recommended safety practices.
- 5. Do not modify equipment unless the manufacturer has approved the modifications.

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#### 7.2.3 Soil/Sediment Sampling Equipment

This section applies to soil sampling equipment including but not limited to hand augers, stainless steel trowels/spoons, bowls, dredges, scoops, split spoons, Macro Core samplers, etc.

- 1. Remove all loose soil from the equipment through manual means.
- 2. Through a combination of scrubbing using soap and water and/or steam cleaning or pressure washing, remove visible dirt/soil from the equipment.
- 3. Rinse the equipment with tap water.

#### **CAUTION**

Do not rinse PE, PVC, and associated tubing with solvents. The appropriate procedures should be defined within the project-specific planning documents. If they are not defined, contact the FOL for guidance. The solvent rinse described in Step 4 may be omitted if groundwater does not contain oil, grease, PAHs, PCBs, or other hard to remove organic materials.

- 4. If the equipment is contaminated or suspected to be contaminated with oil, grease, PAHs, PCBs, or other hard to remove organic materials, rinse the equipment with pesticide-grade isopropanol.
- 5. Rinse the equipment with deionized water.
- 6. To the extent possible, allow components to air dry.
- 7. If the equipment is to be used immediately after decontamination, screen it with a calibrated PID/FID to ensure that all solvents (if they were used) and trace contaminants have been adequately removed.
- 8. After the equipment has dried, wrap it in aluminum foil for storage until use.

Dredges employed in sediment sampling are typically decontaminated as follows:

- Remove the sediment sample from the sampling device
- If sufficient associated surface water is available at the sampling site, place the dredge in the water and flush to remove visible sediment.
- Extract the dredge and wash it in soap and water per the project-specific planning documents.

#### **CAUTION**

When handling dredges, the primary safety concern is trapping fingers or extremities in the larger dredge samplers within the jaws or pinch points of the mechanical jaws. Keep hands, fingers, and extremities away from these pinch and compression points. Either handle the device by the rope or preferably lock the jaws in place to control the potential for closing during maintenance and/or cleaning.

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#### 7.3 Contact Waste/Materials

During the course of field investigations, disposable/single-use equipment becomes contaminated. These items include tubing, trowels, PPE (gloves, overboots, splash suits, etc.), and broken sample containers.

With the exception of the broken glass, single-use articles should be cleaned (washed and rinsed) of visible materials and disposed as normal refuse. The exception to this rule is that extremely soiled materials that cannot be cleaned shall be containerized for disposal in accordance with applicable federal, state, and local regulations.

#### 7.3.1 Investigation-Derived Wastes - Decontamination Wash Waters and Sediments

#### **NOTE**

Requirements for waste storage may differ from one facility to the next. Facility-specific directions for waste storage areas will be provided in project-specific documents, or separate direction will be provided by the Project Manager.

- Assume that all investigation-derived waste (IDW) generated from decontamination activities contains
  the hazardous chemicals associated with the site unless there are analytical or other data to the
  contrary. Waste solution volumes could vary from a few gallons to several hundred gallons in cases
  where large equipment required cleaning.
- 2. Where possible, use filtering systems to extend the use of water within a closed system wash unit to recycle water and to reduce possible waste amounts.

#### NOTE

Containerized waste rinse solutions are best stored in 55-gallon drums (or equivalent containers) that can be sealed until ultimate disposal at an approved facility.

- 3. Label waste storage containers appropriately labeled (see Attachment A).
- 4. Ensure that the IDW storage area is configured to meet the following specifications to permit access to the containers and to conduct spill/leak monitoring, sampling, and extraction when the disposal route is determined:
  - Enclose areas accessible by the general public using construction fencing and signs.
  - Stored materials in 55-gallon drums on pallets with four (or fewer) drums per pallet.
  - Maintain the retaining bolt and label on the outside of storage containers where readily visible.
  - Provide at least 4 feet of room between each row of pallets to allow access to containers for sampling, drum removal, and spill response.
  - As directed in project-specific planning documents, maintain an IDW Inventory List and provide the list to the site Point of Contact at the termination of each shift.
  - Maintain spill response equipment at the IDW storage area in case it is required for immediate access.

Subject	DECONTAMINATION OF FIELD EQUIPMENT	Number SA-7.1	Page 13 of 16
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	<ul> <li>Where possible, use equipment manipulate containers.</li> </ul>	for moving containers. Where no	t possible, obtain help to

Subject DECONTAMINATION OF FIELD EQUIPMENT	Number SA-7.1	Page 14 of 16
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#### **CAUTION**

Each container of water can weigh up to 490 pounds. Each 55-gallon drum of wet soil can weigh more than 750 pounds. Fill drums and temporary containers to 80 percent capacity to minimize spill and handling difficulties. Use drum carts to move filled drums.

See safe lifting techniques provided in Section 4.4 of the Tetra Tech NUS, Inc. Health and Safety Guidance Manual.

When placing drums, keep your fingers out of pinch and smash points such as between the drums. In some cases such as well development and/or purge water, you can place the drums to be filled on the pallet and transport materials in smaller easier to handle containers.

#### 7.4 <u>Decontamination Evaluation</u>

Upon decontamination of equipment, determine the effectiveness of the decontamination process in the following manner:

- Visual evaluation A visual evaluation will be conducted to ensure the removal of particulate matter. This shall be done to ensure that the washing/rinsing process is working as intended.
- Instrument Screening A properly calibrated PID/FID should be used to evaluate the presence of site contaminants and solvents used in the cleaning process. The air intake of the instrument shall be passed over the article to be evaluated. Avoid placing the instrument probe into residual waters. A PID/FID reading greater than the daily established background level requires a repeat of the decontamination process, followed by rescreening with the PID/FID. This sequence must be repeated until no instrument readings greater than the daily established background level are observed. It should be noted that the instrument scan is only viable if the contaminants are detectable within the instrument's capabilities.

#### NOTE

When required by project-specific planning documents, collection of rinsate blanks (see next step) shall be completed without exception unless approval to not collect these samples is obtained from the Project Manager.

- Collection of Rinsate Blanks It is recommended that rinsate samples be collected to:
  - Evaluate the decontamination procedure representing different equipment applications (pumps versus drilling equipment) and different decontamination applications.
  - Single-use disposable equipment The number of samples should represent different types of equipment as well as different lot numbers of single-use articles.
  - The collection and the frequency of collection of rinsate samples are as follows unless specified differently in the project-specific planning documents:
    - Per decontamination method
    - Per disposable article/batch number of disposable articles

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#### NOTE

It is recommended that an initial rinsate sample be collected early in the project to ensure that the decontamination process is functioning properly and to avoid using a contaminated batch of single-use articles. It is recommended that a follow-up sample be collected later during the execution of the project to ensure that those conditions do not change.

Rinsate samples collection may be driven by types of and/or levels of contaminant. Difficult to remove contaminants, oils/greases, some PAHs/PCBs, etc. may also support the collection of additional rinsates due to the obvious challenges to the decontamination process. This is a field consideration to be determined by the FOL.



Subject DECONTAMINATION OF FIELD EQUIPMENT

## STANDARD OPERATING PROCEDURES

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Applicability

Tetra Tech NUS, Inc.

Prepared

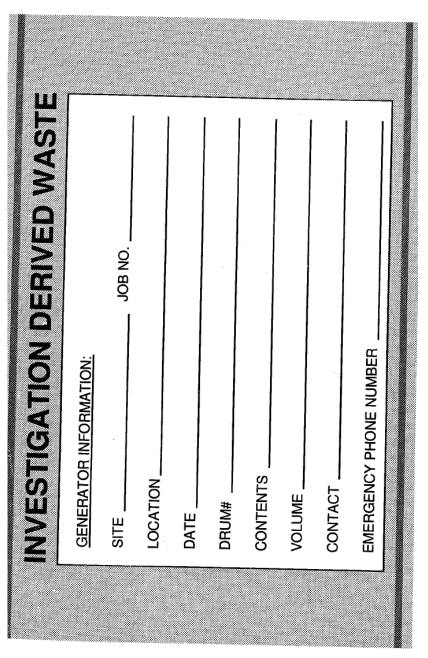
Earth Sciences Department

Approved

Tom Johnston

TE Johnson

## Attachment A iDW Label



# APPENDIX C FIELD DOCUMENTATION FORMS

BORING No.:	TANK FARM 2 CTO WE 30	PROJECT NAME:
DATE:	112G03019	PROJECT NUMBER:
GEOLOGIST:		DRILLING COMPANY:
DRILLER:		DRILLING RIG:

Page \_\_\_ of \_\_\_

						M	ATERIAL DESCRIPTION			PID/FID Readin		FID Reading (ppm)	
Sample No. and Type or RQD	Depth (Ft.) or Run No.	Blows / 6" or RQD (%)	Sample Recovery / Sample Length	ecovery / Change	Soil Density/ Consistency or Rock Hardness	Color	Material Classification	U S C S *	Remarks	Sample	Sampler BZ	Borehole**	Driller BZ**
													1

* When rock coring, enter rock brokenes	5.				
** Include monitor reading in 6 foot interv	als @ borehole. Increase reading frequency if eleva		Drilling Area		
Remarks:				Background (ppm):	
			<u>-</u>		
Converted to Well:	Yes	No	Well I.D. #:		

Æ	TETRA TECH NUS, INC.
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CHAIN OF CUSTODY

NUMBER

4869 | PAGE\_\_\_OF\_\_\_\_

PROJ	ECT NO:		FACILITY:		PROJE	CT MA	NAGER	<u> </u>		PHONE N	JMBER		ı	ABORA	TORY	NAME /	AND CO	NTACT:	
SAMP	LERS (SI	GNATURE)			FIELD	OPERA	TIONS	LEADER		PHONE N	JMBER		. 4	ADDRES	SS				
					CARRI	ER/WA	YBILL N	NUMBER		,			(	CITY, ST	ATE				
STAN RUSH	DARD TA TAT [] hr. []	т 🗆 💮					, ac,			PLAS	ERVA1	or GLA	SS (G)		//				
DATE T	hr		hr.	14 day  OI NOILE	тор бертн (FT)	ВОТТОМ DEPTH (FT)	MATRIX (GW, SO, SW, SD, QC, ETC.)	COLLECTION METHOD GRAB (G) COMP (C)	No. OF CONTAINERS	THE		SIS						CO	MMENTS
						·													
	*** *																		
1 RE	_INQUISH	ED BY			DATE			TIME	1	. RECEIVE	D BY						DA	TE.	TIME
	LINQUISH				DATE	•		TIME		. RECEIVE								TE	TIME
	INQUISH				DATE			TIME		. RECEIVE							DA	TE	TIME
COMI	MENTS				<u> </u>					•									

## Tetra Tech NUS, Inc.

PROJECT:	JOB #:		
LOCATION:	DATE:		
PROJECT MANAGER: F	OL:		
DAILY ACTIVITIES	CHECKLIST		
Startup Checkl	ist		
Activity	Yes	No	N/A
Pertinent site activities/information entered into site logbook			
All onsite personnel listed in logbook			
All onsite personnel listed in logbook  Required medical information onsite for all workers (TtNUS and Sub	contractors)		
Required MSDS's onsite			
Proper equipment calibrations performed (list equipment)			
1			
2			
3			
4			
6 W 4			
Calibration logs filled out Tailgate H&S meeting held prior to beginning field activities			
Required work permits filled out/signed			
Required utility clearances obtained			
Required PPE onsite and in use			
Information required to be posted is in place			
(OSHA poster, hospital route, key phone numbers, etc.)			
Exit Checklis	4		
LAR OHECKIS			
Activity	Yes	No	N/A
Logbooks completely and comprehensively filled out			<u></u>
Field forms complete and accounted for/properly filed			
Samples properly packaged/shipped			
COCS taxed to appropriate in-nouse personnel			
All equipment accounted for, on charge if needed, and properly sec	ured		
All personnel accounted for			
All personnel accounted for Arrangements made for upcoming work (permits, clearances, equip	ment, etc.)		<u> </u>
Site properly secured			
•			

Note - not all items listed apply to every job, and some additional requirements may apply on a job-specific basis.

TETRA TECH	I NUS, INC.	FIELD INSTRUMENT CALIBRATION LOG							
INSTRUMENT NAME:		MODEL No.:							
CALIBRATION DATE	INITIAL READING	PROCEDURE	FINAL READING	SIGNATURE	COMMENTS				



## TETRA TECH NUS, INC. FIELD MODIFICATION RECORD

Site Name: _			Location:	
Project Number	er:		Task Assignment:	
To:	Loc	ation:	D	ate:
Description:				
Reason for Ch	nange:			
D	J. A. et'a.			
Recommende	d Action:			
	ns Leader (Signature):			Date:
Disposition/Ac	tion:			
	(0: )			<b>.</b>
Project Manag	ger (Signature):			Date:
Distribution:	Program Manager:			Others as Required:
	Project Manager:			
	Quality Assurance Officer: Field Operations Leader:			
	Project File:			- -



# TETRA TECH NUS FIELD TASK MODIFICATION REQUEST FORM

Project/Installation Name	CTO & Project Number	Task Mod. Number
Modification To (e.g. Work Plan)	Site/Sample Location	Date
Activity Description:		
Reason for Change:		
Recommended Disposition:		
Field Operations Leader (Signature	<del>)</del>	Date
	<del>&gt;</del> )	Date
	<del>=</del> )	Date
		Date
	<b>⇒</b> )	Date
Field Operations Leader (Signature Approved Disposition:  Project/Task Order Manager (Signature)		Date
Approved Disposition:		



### PHOTOIONIZATION DETECTOR FIELD CALIBRATION LOG

Serial No.: Model No.:				Decal No.:		
Site Name/Location: _Tank Farm 2 - NAVSTA Newport Newport, RI				etra Tech NUS Charge No.: <u>112</u>	2G03019 / CTO WE30	
CALIBRATION DATE	STANDARD GAS- ISOBUTYLENE	(AM) CALIBRATION READING Isobutylene Equiv. (ppm)	(PM) CALIBRATION CHECK Isobutylene Equiv. (ppm)	SIGNATURE	COMMENTS	

CALIBRATION DATE	STANDARD GAS- ISOBUTYLENE	CALIBRATION READING Isobutylene Equiv. (ppm)	CALIBRATION CHECK Isobutylene Equiv. (ppm)	SIGNATURE	COMMENTS
	Lot # ppm	/		AM: PM:	
	Lot # ppm	/		AM: PM:	
	Lot # ppm	/		AM: PM:	
	Lot # ppm	/		AM: PM:	
	Lot # ppm	/		AM: PM:	
	Lot # ppm	/		AM: PM:	
	Lot # ppm	/		AM: PM:	
	Lot # ppm	/		AM: PM:	
	Lot # ppm	/		AM: PM:	



			Page_	of
Project Number: 112 Sample Location: QA Sample Type:	IK FARM 2 G03019 o Blank	Sample ID Number: Sampled By: C.O.C. Number:		
[ ] Sou (Fiel	urce Water Blank d Blank)	[ ] Other Blank		
SAMPLING DATA:		WATER SOURCE:		9 9 99
Date: Time: Method: Direct Pour		[ ] Laboratory Prepared [ ] Purchased [ ] Other	[]Tap []Fire Hy	drant
PURCHASED WATER INFORM (If Applicable as Source or		RINSATE INFORMATIO (if Applicable):	N	
Product Name: Reagent Grade	Water (DIUF)	Media Type:		
Supplier:		Equipment Used:		
Manufacturer:	_	Equipment Type:		
Order Number:			Dedicated	
Lot Number:		[	[ ] Reusable	
Expiration Date:		[	] Disposable	
SAMPLE COLLECTION INFOR	MATION:			
SAMPLE COLLECTION INFOR	MATION:  Preservative	Container Req	uirements	Collected
		Container Req	uirements	Collected YES / NO
Analysis		Container Req	uirements	YES/NO YES/NO
<b>Analysis</b> VOC		Container Req	uirements	YES/NO YES/NO YES/NO
Analysis VOC GRO PAH ExTPH		Container Req	uirements	YES/NO YES/NO YES/NO YES/NO
Analysis VOC GRO PAH		Container Req	uirements	YES/NO YES/NO YES/NO YES/NO YES/NO
Analysis VOC GRO PAH ExTPH		Container Req	uirements	YES/NO YES/NO YES/NO YES/NO
Analysis VOC GRO PAH ExTPH Metals		Container Req	uirements	YES/NO YES/NO YES/NO YES/NO YES/NO

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#### TETRA TECH NUS, INC.

### RECORD OF FIELD WORK ORIENTATION

SITE:				JO	B No.:	<del>,</del>	
WORK ASSIG	GNMENT No.: _						
TASK OR AC	CTIVITY:			DATE	OF ORIENTA	ΓΙΟΝ:	
PERSONNEL	. ATTENDING	<u>TI</u>	RAINERS	FOL	:	PROJECT MO	GR.
1.		1.		1.	1.		
2.		2.					
3.		3.					
4.							
5.							
6.							
7.							
VERIFICATIO	ONS (CHECK A	ND INITIAL B	Y ATTENDEE	ES)			
WORK PLAN REVIEWED			ITE/EQUIP. SECURITY REVIEWED	EQUIPMENT OPERATION		PURCHASING	INITIALS
1							
2							
3							
6 7.							
	RE	TURN ORIGIN	NAL TO THE	QUALITY ASSU	RANCE OFFIC	ER	
Copies to:	PROJECT FIL	.E:					
	PROJECT MAI	NAGER:					
	PROGRAM MA	ANAGER:				_	

Tt.	TETRA TECH NUS, INC.

### SAMPLE COLLECTION SUMMARY RECORD

	TETRA TECH NUS CHARGE NUMBER:									
SAMPLING EVENT:				C/	ASE	۱O.: _			DAS NO.:	
DATE	TIME	SAMPLE LOCATION	FIELD QC							COMMENTS

TETRA TECH NUS, INC.
Site Name: Tank Farm 2 NAVSTA

### SITE ENTRY LOG

Site Name: Tank Farm 2 NAVSTA Newport	Date: .
Location: Newport, RI	Project Number: 112G03019 / CTO WE 30

NAME	REPRESENTING	TIME IN (HOURS)	TIME OUT (HOURS)	INITIALS
	TTNUS			
	TTNUS			

TtNUS Form 0002



#### SOIL AND SEDIMENT SAMPLE LOG SHEET

Page<sub></sub> of Project Site Name: Tank Farm 2 NAVSTA Newport, Newport, RI Sample ID No.: Project No.: 112G03019. / CTO WE30 Sample Location: Sampled By: 1 Surface Soil C.O.C. No.: Subsurface Soil ] Sediment Type of Sample: ] Low Concentration Other: ] High Concentration [ ] QA Sample Type: GRAB SAMPLE DATA: VOC: Depth Interval Description (Sand, Silt, Clay, Moisture, etc.) Date: Color Time: Method: Monitor Reading (ppm): COMPOSITE:SAMPLE:DATA: ALL:NON-YOC Date: Description (Sand, Silt, Clay, Moisture, etc.) Time **Depth Interval** Color Method: Monitor Readings (Range in ppm): SAMPLE COLLECTION:INFORMATION: Analysis Preservative **Container Requirements** Collected Other MeOH, 4°C 1 x 40 mL vial YES / NO VOC YES / NO NaHSO4, 4°C 2 x 40 mL vial 1 x 40 mL vial GRO MeOH, 4°C YES / NO percent moisture 4°C YES / NO one 2 oz jar DI, 4°C PAH 2 x 40 mL vial YES / NO XTPH one 8 oz jar YES / NO one 4 oz jar YES / NO Dioxins 4°C YES / NO Metals 4°C one 4 oz jar OBSERVATIONS 7 NOTES: Circle if Applicable: Signature(s): **Duplicate ID No.:** MS/MSD

# APPENDIX D PROJECT-SPECIFIC FIELD TASK PROCEDURES

#### **APPENDIX D-1**

#### MOBILIZATION/DEMOBILIZATION AND UTILITY CLEARANCE

Mobilization includes procurement of field equipment, supplies, and subcontractors; mobilization of personnel, subcontractors, equipment, and supplies to the field; coordination with Dig-safe and the NAVSTA Environmental Office; a site walkover; and field orientation meetings to prepare for field work. Mobilization will be coordinated with the NAVSTA Environmental Office a minimum of 1 week in advance of the initiation of field activities.

TtNUS will be responsible for obtaining clearance of all underground utilities at all sampling locations at least 72 hours prior to beginning on-site work. TtNUS will conduct a public and Navy Dig-safe clearance of the Site. TtNUS will coordinate with local utility contractors and the NAVSTA Environmental Office, marking drilling locations on the ground with stakes, pin flags, and white spray paint, visiting the Site to meet with NAVSTA Environmental Office utility marking crews, and making adjustments as needed (to avoid overhead hazards, utility lines, etc). Utility clearance will be performed in accordance with TtNUS SOP HS-1.0 (Appendix A).

A field orientation meeting will be held prior to beginning the field program. The purpose of this meeting is to review the scope of the field study including site description, objectives of the field investigation, sampling locations, sampling methods, field QC samples, health and safety requirements, NAVSTA protocol and chain-of-command. The field orientation meeting will be attended by the field staff, project manager, lead chemist, and health and safety officer. Field team members will review this SAP, applicable SOPs (Appendix A), and applicable Field Documentation Forms (Appendix B) and they will document that they have read the SAP by signing Worksheet #4. All subcontractors working at the Site will be provided with the site-specific health and safety plan, as well as their specifications and scopes of work.

Portable sanitary facilities, vehicles, and supply storage containers will be procured for the Site. Drillers will bring a drill rig, support truck, and a water trailer to the Site for decontamination purposes. Drums (55-gallon) will be brought to the site to hold drill cuttings. It is assumed that it will be acceptable to store drums outside and not require a storage box at this site.

Demobilization includes removing field equipment and supplies from the Site, returning rented equipment, managing IDW, performing general site cleanup, organizing and finalizing field paperwork, and entering field records/data into the Site database.

#### **APPENDIX D-2**

#### PROJECT-SPECIFIC SAMPLING PROCEDURES

Soil sampling at Tank Farm 2 will be performed according to the SOPs and project-specific sampling procedures described below. The SOPs and field documents referred to below are included in Appendix B and C, respectively.

Quality control (QC) samples will be collected as part of the investigation, including field duplicates, rinsate blanks, field blanks, and trip blanks. Samples will also be assigned on the chain-of-custody (COC) form for laboratory QC analyses. Worksheet #20 summarizes the QC samples to be collected for each matrix.

The sample locations are presented on Figures 5 through 9 and on Worksheet #18. Worksheet #18 also presents the analytical groups for each sample location. Worksheet #19 presents the analytical methods, sample container types, preservative requirements, and the maximum holding times for the associated analyses.

Samples will be identified in accordance with the sample location identification system presented in Worksheet #27. All laboratory analytical samples will be kept on ice in coolers and will be shipped with appropriate QC samples. Samples will be handled and delivered in accordance with the COC procedures detailed in Worksheet #27.

#### 1.0 SOIL SAMPLING PROCEDURES

Soil borings will be advanced at Tank Farm 2 at locations for continuous soil sampling. A variety of drilling techniques may be used depending on the conditions encountered at the site. Direct-push drilling, hand augers and/ or other hand tools may be used.

Drilling will be performed by a subcontractor according to the procedures in SOP SA-2.5 for DPT or SOP GH-1.3 for other drilling methods. All down-hole drilling equipment will be steam-cleaned before use at each boring. For direct-push, a 2-inch ID, 4 or 5-foot long, stainless steel, macro-core sampler with an inner acetate liner will be used to collect the soil samples. All required information will be recorded on the boring log, in according with SOP GH-1.5.

Soil samples will be collected for laboratory analysis from the soil cores in accordance with SOP SA-1.3 and the procedures described below. Please refer to Worksheet # 18 for details on sample interval selection in Category 1 and Category 2 areas. The boring log will act as the sample log sheet for samples collected.

For each interval to be sampled for GRO, the split-barrel sampler or acetate liner will be opened, visually inspected, and scanned with a photo-ionization detector (PID) portable monitoring instrument. The GRO aliquots will be collected from the most heavily contaminated portion of the split-barrel sampler, based on the initial field screening results and/or visual and olfactory observations. The specific depth of the GRO samples within the 2-foot split barrel sample will be recorded. The remainder of the soil from the sample interval will be uniformly hand mixed to form a composite sample and split into aliquots for the remaining non-volatile analyses (PAHs, metals, and (at select depths) dioxins; or PCB analysis, see Worksheet #18 for details). For these aliquots, the sampling procedures below for non-volatile parameters will be followed.

#### Soil Sampling Procedures for Volatile Laboratory Samples (Grab)

Each soil sample for volatile analysis (GRO) will be collected using a dedicated cut syringe or equivalent device. The GRO aliquot will be placed in methanol-preserved vials with a septa cap. The vials will be maintained at  $\leq$  6 °C for up to 14 days. The following procedures should be followed for the soil volatile sample collection:

- 1. Label a bag containing two 40-mL amber vials containing 5 mL of methanol for the GRO aliquots, with the sample location number and a bottle identifier such as A, B, etc.
- 2. Collect approximately 5 grams of sample by coring or stabbing the soil with a dedicated 10-mL pre-cut syringe. Extrude the sample into one of the 40-mL VOC vials containing 5 mL of preservative (methanol). The soil must be immersed in the preservative; recollect the sample using a smaller volume if necessary. Avoid touching the threads on the vial's neck and avoid loss of preservative by evaporation. Cap the vial and invert it several times to mix the sample.

3. Weigh each sample vial to the nearest 0.01 gram and record the weight on the field log sheet if samples are being shipped. Repeat the sample collection procedure for the remaining vials. Pack and ship to the laboratory. Include the field log sheet containing the sample weight information with the samples.

Quality assurance and quality control samples will also be collected, see Worksheet #20 for details. Following the collection of the first set of volatile containers, collect the field duplicate from the same sampling interval.

Every effort should be made to obtain the percent moisture soil aliquot (see below) as close as possible to the location where the volatile sample was collected.

#### Soil Sample for Percent Moisture

Fill one 2-oz container with soil representing the same locations where the 40-mL volatile vial samples were collected. Every effort should be made to obtain the percent moisture soil aliquot as close as possible to the location where the volatile samples were collected. This is a grab sample (not composited).

#### Soil Sampling Procedures for Non-GRO Parameters

- Record all required data on the boring log, which will also serve as the soil sample log sheet. Include the sampling equipment, sampling personnel, date, time, depth of sample, and sample analyses. Use the boring log to record soil descriptions, depth of strata changes, and sample depth intervals. Classify the soil sample visually using the Unified Soil Classification System (ASTM D-2488-98).
- 2. Label appropriate sample jars with the sample location number, sampler's name, date, and analytical fractions.
- 3. Transfer the soil from the sampler to a decontaminated stainless-steel bowl using only decontaminated stainless steel trowels, and homogenize the sample.

- 4. If there is insufficient sample volume to fill all the containers for analyses, an equal amount of material from the intervals immediately above and below the selected sample interval may be used to supplement the composite sample to ensure sufficient sample quantity for all analyses. Alternatively, a second boring immediately adjacent to the original boring could be advanced to the desired depth to obtain additional sample volume. Document the method used in the boring log.
- 5. Remove any large particles such as twigs, gravel or artificial fill too large to be sent for analysis. Document the removal of material on the boring log.
- 6. Fill the appropriate sample containers with the soil sample.
- 7. For field duplicate samples, after homogenization, fill one set of sample containers for the original sample and fill another set for the field duplicate sample.
- 8. Ensure that the samples are properly labeled, maintained in coolers with ice, and that the COC procedures described in Worksheet #27 are followed. Package and ship the sample coolers to the appropriate laboratory for overnight delivery.
- 9. Decontaminate the sampling equipment before reuse.

Care should be taken in handling all soil samples to ensure that the exterior of the sample containers are clean and free of soils before shipping to the laboratory.

#### **APPENDIX D-3**

#### INVESTIGATION DERIVED WASTE MANAGEMENT

It is anticipated that waste materials will be generated during the field investigation. These materials include:

- Decontamination fluid;
- Used personal protective equipment (PPE);
- Used sampling equipment;
- Drill cuttings and excess soil samples;

Investigation-derived waste will be managed as described below, in accordance with RIDEM regulations (Rules and Regulations for Hazardous Waste Management, DEM OWM-HW01-07).

Visibly clear phosphate-free detergent wash water and rinse water decontamination fluids from sampling equipment will be released to the ground upon rinsing, in the immediate vicinity of its point of generation. The decontamination rinse water will be contained in 55-gallon drums or bulk containers.

Used PPE, such as sampling gloves, Tyvek coveralls, paper towels, or other materials will be bagged and sealed prior to disposal as general refuse. If PPE becomes grossly contaminated, it will be segregated from other PPE, labeled and staged as "contaminated material." Contaminated material will be drummed and staged in the IDW area designated by the Navy. TtNUS will arrange for off-site disposal of drums by a licensed waste hauler at an approved facility.

Used disposable sampling equipment, which generally has minor contamination, will be disposed of with the PPE as general refuse. Grossly contaminated disposable equipment, as determined at the discretion of the sampler, will require segregation from other equipment and proper disposal.

Drill cuttings and excess soil samples will be contained in 55-gallon drums or bulk containers and staged at a secure area. Any drums used for storage will be clearly marked with a grease pencil or other water-resistant marker to indicate the borehole from which the cuttings were removed. The word "soil" will be used to differentiate between drummed cuttings and drummed purge water or development water. The drums or bulk containers will be staged in an orderly fashion, with

proper spacing, in an area designated by the Navy. Composite samples will be collected from the filled drums to characterize the waste generated. Data from the analysis of these composite samples will be used for characterization of the materials, and, based on these analyses, the material will be manifested for shipment off-site to a disposal facility. Waste pickup and disposal will be conducted in coordination with the NAVSTA waste management office, who takes ownership of the waste after characterization.

TtNUS will be responsible for arranging the removal and proper disposal of all accumulated waste materials following completion of the investigation. Manifests and shipping papers will be signed by a representative of the NAVSTA waste management office. Disposal will be arranged with licensed waste haulers at approved receiving facilities. Characterization analyses will be conducted by the waste disposal subcontractor.

### APPENDIX D-4 LAND SURVEYING

After completion of sample collection, the coordinates of soil borings, and other pertinent features will be determined by a Rhode Island registered land surveyor. The coordinates of the features will be incorporated into the NAVSTA Newport GIS database and used for site mapping.

Surveying activities will establish the horizontal coordinates of borings. The survey will also establish ground surface elevations at these investigation locations.

All vertical measurements will be surveyed in United States Geological Survey (USGS) National Geodetic Vertical Datum of 1929 (NGVD 1929) coordinates, in feet. All horizontal measurements will be in Rhode Island State Plane coordinates, using 1983 North American Datum (NAD 1983), in feet.

Horizontal locations will be measured to the nearest 0.1 foot NAD 1983. Vertical positions of well risers will be measured to the nearest 0.01 foot NVGD 1929. Vertical and horizontal control will be brought to the Site using local benchmark information and temporary benchmarks will be established as needed.

The subcontractor deliverable will include a base map of the Site, showing existing permanent features as well as sample stations (borings, wells, and other stations of interest). General topography will be presented with five-foot contours.

## APPENDIX E ANALYTICAL SPECIFICATIONS

# ATTACHMENT A - TABLE A-1 SOIL PROJECT-REQUIRED TARGET ANALYTES AND PSLs CTO WE59, TANK FARM 3 NAVSTA NEWPORT, NEWPORT, RI REMEDIAL INVESTIGATION, CHEMICAL ANALYSES

Matrix: Soil

	CAS	Soil PSL	LOQ	Ka	tahdin Lin	nits
Analyte	Number	(mg/kg)	Goal	LOQ	LOD	DL
	Number	(ilig/kg)	(mg/kg)	(mg/kg)	(mg/kg)	(mg/kg)
VOCs by SW-846 8260B						
1,2,4-Trimethylbenzene	95-63-6	6.2	2.1			
1,2-Dibromoethane (EDB)	106-93-4	0.0001	0.000033			
1,3,5-Trimethylbenzene	108-67-8	7.2	2.4			
2-Butanone (MEK)	78-93-3	89.6	30			
2-Hexanone	591-78-6	12.6	4.2			
4-Methyl-2-pentanone						
(MIBK)	108-10-1	443	150			
Acetone	67-64-1	2.5	0.83			
Benzene	71-43-2	0.2	0.67			
Bromoform	75-25-2	15.9	5.3			
Bromomethane	74-83-9	0.235	0.078			
Carbon disulfide	75-15-0	0.0941	0.031			
Cyclohexane	110-82-7	700	230			
Ethylbenzene	100-41-4	5.16	1.7			
Isopropylbenzene	98-82-8	27	9			
m,p-Xylenes <sup>4</sup>	179601-23-1	95	32			
Methyl acetate	79-20-9	7800	2600			
Methylcyclohexane	108-87-2	490	160			
Methyl-tert-butyl ether	1634-04-4	43	1.3			
Naphthalene	91-20-3	0.8	0.27			
n-Butylbenzene	104-51-8					
n-Propylbenzene	103-65-1	340	11			
o-Xylene⁴	95-47-6	95	32			
p-Isopropyltoluene	99-87-6					
sec-Butylbenzene	135-98-8					
Styrene	100-42-5	2.9	0.97			
tert-Butylbenzene	98-06-6					
Toluene	108-88-3	5.45	1.8			
Xylenes (total)	1330-20-7	63	21			

	CAS	Soil PSL	LOQ	Ka	atahdin Lin	nits
PAHs by SW-846 8270D SIM	Number	(mg/kg)	Goal (mg/kg)	LOQ (mg/kg)	LOD (mg/kg)	DL (mg/kg)
2-Methylnaphthalene	91-57-6	29	9.7			
Acenaphthene	83-32-9	20	6.7			
Acenaphthylene	208-96-8	23	7.7			
Anthracene	120-12-7	29	9.7			
Benzo(a)anthracene	56-55-3	0.15	0.05			
Benzo(a)pyrene	50-32-8	0.015	0.005			
Benzo(b)fluoranthene	205-99-2	0.15	0.05			
Benzo(g,h,i)perylene	191-24-2	0.8	0.27			
Benzo(k)fluoranthene	207-08-9	0.9	0.3			
Chrysene	218-01-9	0.4	0.13			
Dibenzo(a,h)anthracene	53-70-3	0.015	0.005			
Fluoranthene	206-44-0	20	6.7			
Fluorene	86-73-7	28	9.3			
Indeno(1,2,3-c,d)pyrene	193-39-5	0.15	0.05			
Naphthalene	91-20-3	0.8	0.27			
Phenanthrene	85-01-8	29	9.7			
Pyrene	129-00-0	1.1	0.37			
PCBs by SW-846 8082A						
Aroclor-1016	12674-11-2	0.39	0.13			
Aroclor-1221	11104-28-2	0.14	0.047			
Aroclor-1232	11141-16-5	0.14	0.047			
Aroclor-1242	53469-21-9	0.22	0.073			
Aroclor-1248	12672-29-6	0.22	0.073		_	
Aroclor-1254	11097-69-1	0.11	0.037			
Aroclor-1260	11096-82-5	0.22	0.073			
Aroclor-1262	37324-23-5	0.22	0.073			
Aroclor-1268	11100-14-4	0.22	0.073			

	CAS	Call DCI	LOQ	Ka	atahdin Lim	nits
Metals by SW-846 6020A/7471B	CAS Number	Soil PSL (mg/kg)	Goal (mg/kg)	LOQ (mg/kg)	LOD (mg/kg)	DL (mg/kg)
Aluminum	7429-90-5	50	17			
Antimony	7440-36-0	0.27	0.09			
Arsenic	7440-38-2	0.39	0.13			
Barium	7440-39-3	330	110			
Beryllium	7440-41-7	1.5	0.5			
Cadmium	7440-43-9	0.36	0.12			
Calcium	7440-70-2					
Chromium	7440-47-3	0.29	0.097			
Cobalt	7440-48-4	2.3	0.77			
Copper	7440-50-8	28	9.3			
Iron <sup>8</sup>	7439-89-6	200	67			
Lead	7439-92-1	11	3.7			
Magnesium	7439-95-4					
Manganese	7439-96-5	180	60			
Mercury	7439-97-6	0.1	0.033			
Nickel	7440-02-0	38	13			
Potassium	7440-09-7					
Selenium	7782-49-2	0.52	0.17			
Silver	7440-22-4	4.2	1.4			
Sodium	7440-23-5					
Thallium	7440-28-0	0.0569	0.019			
Vanadium	7440-62-2	2	0.67			
Zinc	7440-66-6	46	15			
Petroleum Hydrocarbons						
GRO (C5-C12)						
ExTPH (C8-C44)						
TPH		500	170			

#### ATTACHMENT A - TABLE A-2 SOIL PROJECT-REQUIRED TARGET ANALYTES AND PSLs CTO WE30, TANK FARM 2 NAVSTA NEWPORT, NEWPORT, RI

#### REMEDIAL INVESTIGATION, CHEMICAL ANALYSES

Matrix: Soil

Matrix: Soil	CAS	Call DCI	LOQ	Ka	tahdin Lim	its
Analyte	Number	Soil PSL (mg/kg)	Goal (mg/kg)	LOQ (mg/kg)	LOD (mg/kg)	DL (mg/kg)
PAHs by SW-846 8270 SIM					, -	
2-Methylnaphthalene	91-57-6	29	9.7			
Acenaphthene	83-32-9	20	6.7			
Acenaphthylene	208-96-8	23	7.7			
Anthracene	120-12-7	29	9.7			
Benzo(a)anthracene	56-55-3	0.15	0.05			
Benzo(a)pyrene	50-32-8	0.015	0.005			
Benzo(b)fluoranthene	205-99-2	0.15	0.05			
Benzo(g,h,i)perylene	191-24-2	0.8	0.27			
Benzo(k)fluoranthene	207-08-9	0.9	0.3			
Chrysene	218-01-9	0.4	0.13			
Dibenzo(a,h)anthracene	53-70-3	0.015	0.005			
Fluoranthene	206-44-0	20	6.7			
Fluorene	86-73-7	28	9.3			
Indeno(1,2,3-c,d)pyrene	193-39-5	0.15	0.05			
Naphthalene	91-20-3	3.6	1.2			
Phenanthrene	85-01-8	29	9.7			
Pyrene	129-00-0	1.1	0.37			
Metals by SW-846 6020A/7471B						
Aluminum	7429-90-5	50	17			
Antimony	7440-36-0	0.27	0.09			
Arsenic	7440-38-2	0.39	0.13			
Barium	7440-39-3	330	110			
Beryllium	7440-41-7	10	3.3			
Cadmium	7440-43-9	0.36	0.12			
Calcium	7440-70-2					
Chromium	7440-47-3	0.29	0.097			
Cobalt	7440-48-4	2.3	0.77			
Copper	7440-50-8	28	9.3			
Iron	7439-89-6	200	66.7			
Lead	7439-92-1	11	3.7			
Magnesium	7439-95-4					
Manganese	7439-96-5	180	60			
Mercury	7439-97-6	0.1	0.033			
Nickel	7440-02-0	38	13			
Potassium	7440-09-7					
Selenium	7782-49-2	0.52	0.17			
Silver	7440-22-4	4.2	1.4			
Sodium	7440-23-5					
Thallium	7440-28-0	0.0569	0.019			

#### Matrix: Soil

	CAS	Soil PSL	LOQ	Ka	tahdin Lim	its
Analyte	Number	(mg/kg)	Goal (mg/kg)	LOQ (mg/kg)	LOD (mg/kg)	DL (mg/kg)
Vanadium	7440-62-2	2	0.67			
Zinc	7440-66-6	46	15			
PCBs by SW-846 8082A						
Aroclor 1016	12674-11-2	0.39	0.13			
Aroclor 1221	11104-28-2	0.14	0.047			
Aroclor 1232	11141-16-5	0.14	0.047			
Aroclor 1242	53469-21-9	0.22	0.073			
Aroclor 1248	12672-29-6	0.22	0.073			
Aroclor 1254	11097-69-1	0.11	0.037			
Aroclor 1260	11096-82-5	0.22	0.073			
Petroleum						
Hydrocarbons						
GRO (C5-C12)						
ExTPH (C8-C44)						
TPH		500	170			

## ATTACHMENT A - TABLE A-3 GROUNDWATER PROJECT-REQUIRED TARGET ANALYTES AND PSLs CTO WE59, TANK FARM 3 NAVSTA NEWPORT, NEWPORT, RI REMEDIAL INVESTIGATION, CHEMICAL ANALYSES

**Matrix: Groundwater** 

watrix: Groundwater		Ground	Project	Kat	ahdin Lir	nits
Analyte	CAS Number	water PSL (µg/L)	LOQ Goal (µg/L)	LOQ (µg/L)	LOD (µg/L)	DL (µg/L)
1,2-Dibromoethane (EDB) by SW-846 8011	106-93-4	0.0065	0.0022			
VOCs by SW-846 8260B						
1,2,4-Trimethylbenzene	95-63-6	1.5	0.5			
1,3,5-Trimethylbenzene	108-67-8	8.7	2.9			
2-Butanone (MEK)	78-93-3	490	160			
2-Hexanone	591-78-6	3.4	1.1			
4-Methyl-2-pentanone (MIBK)	108-10-1	100	67			
Acetone	67-64-1	1200	400			
Benzene	71-43-2	0.39	0.13			
Bromoform	75-25-2	7.9	0.0028			
Bromomethane	74-83-9	0.7	0.23			
Carbon disulfide	75-15-0	72	24			
Cyclohexane	110-82-7	100	33			
Ethylbenzene	100-41-4	1.3	0.5			
Isopropylbenzene	98-82-8	39	13			
	179601-23-					
m,p-Xylenes	1	19	6.3			
Methyl acetate	79-20-9	1600	533			
Methylcyclohexane	108-87-2					
Methyl-tert-butyl ether	1634-04-4	12	4			
Naphthalene	91-20-3	0.14	0.047			
n-Butylbenzene	104-51-8	78	26			
n-Propylbenzene	103-65-1	53	18			
o-Xylene	95-47-6	19	6.3			
p-Isopropyltoluene	99-87-6					
sec-Butylbenzene	135-98-8	78	26			
Styrene	100-42-5	110	37			
tert-Butylbenzene	98-06-6	78	26			
Toluene	108-88-3	86	29			
Xylenes (total)	1330-20-7	19	6.3			
PAHs by SW-846 8270D SIM						
2-Methylnaphthalene	91-57-6	2.7	0.9			
Acenaphthene	83-32-9	40	13			
Acenaphthylene	208-96-8	40	13			
Anthracene	120-12-7	130	43			
Benzo(a)anthracene	56-55-3	0.029	0.0097			
Benzo(a)pyrene	50-32-8	0.0029	0.00097			

**Matrix: Groundwater** 

Matrix: Groundwater		Ground	Project	Kat	ahdin Lir	nits
Analyte	CAS Number	water PSL (µg/L)	LOQ Goal (µg/L)	LOQ (µg/L)	LOD (µg/L)	DL (µg/L)
Benzo(b)fluoranthene	205-99-2	0.029	0.0097			
Benzo(g,h,i)perylene	191-24-2	8.7	2.9			
Benzo(k)fluoranthene	207-08-9	0.29	0.097			
Chrysene	218-01-9	2.9	0.97			
Dibenzo(a,h)anthracene	53-70-3	0.0029	0.00097			
Fluoranthene	206-44-0	63	21			
Fluorene	86-73-7	22	7.3			
Indeno(1,2,3-c,d)pyrene	193-39-5	0.029	0.0097			
Naphthalene	91-20-3	0.14	0.047			
Phenanthrene	85-01-8	8.7	2.9			
Pyrene	129-00-0	8.7	2.9			
PCBs by SW-846 8082A						
Aroclor-1016	12674-11-2	0.11	0.034			
Aroclor-1221	11104-28-2	0.004	0.00013			
Aroclor-1232	11141-16-5	0.004	0.00013			
Aroclor-1242	53469-21-9	0.034	0.011			
Aroclor-1248	12672-29-6	0.034	0.011			
Aroclor-1254	11097-69-1	0.034	0.011			
Aroclor-1260	11096-82-5	0.034	0.011			
Aroclor-1262	37324-23-5	0.034	0.011			
Aroclor-1268	11100-14-4	0.034	0.011			
Metals by SW-846 6020A/74						
Aluminum	7429-90-5	1600	533			
Antimony	7440-36-0	0.6	0.2			
Arsenic	7440-38-2	0.045	0.015			
Barium	7440-39-3	290	97			
Beryllium	7440-41-7	4	1.3			
Cadmium	7440-43-9	0.69	0.23			
Calcium	7440-70-2					
Chromium	7440-47-3	0.031	0.010			
Cobalt	7440-48-4	0.47	0.16			
Copper	7440-50-8	62	21			
Iron	7439-89-6	1100	367			
Lead	7439-92-1	15	5			
Magnesium	7439-95-4					
Manganese	7439-96-5	32	11			
Mercury	7439-97-6	0.067	0.022			
Nickel	7440-02-0	30	10			
Potassium	7440-09-7					
Selenium	7782-49-2	7.8	2.6			
Silver	7440-22-4	7.1	2.4			
Sodium	7440-23-5					
Thallium	7440-28-0	2	0.67			
Vanadium	7440-62-2	7.8	2.6			
Zinc	7440-66-6	470	157			

## ATTACHMENT A - TABLE A-4 SEDIMENT PROJECT-REQUIRED TARGET ANALYTES AND PSLs CTO WE59, TANK FARM 3 NAVSTA NEWPORT, NEWPORT, RI REMEDIAL INVESTIGATION, CHEMICAL ANALYSES

**Matrix: Sediment** 

watrix: Sediment			Project	Ka	tahdin Lin	nits
Analyte	CAS Number	Sed PSL (mg/kg)	LOQ Goal (mg/kg)	LOQ (mg/kg)	LOD (mg/kg)	DL (mg/kg)
VOCs by SW-846 8260B						
1,2,4-Trimethylbenzene	95-63-6	6.2	2.1			
1,2-Dibromoethane (EDB)	106-93-4	0.034	0.011			
1,3,5-Trimethylbenzene	108-67-8	78	26			
2-Butanone (MEK)	78-93-3	0.27	0.09			
2-Hexanone	591-78-6	0.022	0.0073			
4-Methyl-2-pentanone (MIBK)	108-10-1	0.033	0.011			
Acetone	67-64-1	0.0087	0.0029			
Benzene	71-43-2	0.16	0.053			
Bromoform	75-25-2	0.654	0.22			
Bromomethane	74-83-9	0.73	0.24			
Carbon disulfide	75-15-0	0.000851	0.00028			
Cyclohexane	110-82-7	700	230			
Ethylbenzene	100-41-4	1.1	0.37			
Isopropylbenzene	98-82-8	0.08	0.027			
	179601-					
m,p-Xylenes	23-1	59	20			
Methyl acetate	79-20-9	7800	2600			
Methylcyclohexane	108-87-2					
Methyl-tert-butyl ether	1634-04-4	43	14			
Naphthalene	91-20-3	0.176	0.059			
n-Butylbenzene	104-51-8					
n-Propylbenzene	103-65-1	340	110			
o-Xylene	95-47-6	69	23			
p-Isopropyltoluene	99-87-6					
sec-Butylbenzene	135-98-8					
Styrene	100-42-5	0.559	0.19			
tert-Butylbenzene	98-06-6					
Toluene	108-88-3	0.05	0.017			
Xylenes (total)	1330-20-7	0.16	0.053			
PAHs by SW-846 8270D SIM						
2-Methylnaphthalene	91-57-6	0.0202	0.0067			
Acenaphthene	83-32-9	0.0067	0.0022			
Acenaphthylene	208-96-8	0.0059	0.002			
Anthracene	120-12-7	0.0572	0.019			
Benzo(a)anthracene	56-55-3	0.108	0.036			
Benzo(a)pyrene	50-32-8	0.015	0.005			

#### **Matrix: Sediment**

watrix. Sediment			Project	Ka	tahdin Lin	nits
Analyte	CAS Number	Sed PSL (mg/kg)	LOQ Goal (mg/kg)	LOQ (mg/kg)	LOD (mg/kg)	DL (mg/kg)
Benzo(b)fluoranthene	205-99-2	0.13	0.043			
Benzo(g,h,i)perylene	191-24-2	0.17	0.057			
Benzo(k)fluoranthene	207-08-9	0.24	0.08			
Chrysene	218-01-9	0.166	0.055			
Dibenzo(a,h)anthracene	53-70-3	0.015	0.005			
Fluoranthene	206-44-0	0.423	0.14			
Fluorene	86-73-7	0.0774	0.026			
Indeno(1,2,3-c,d)pyrene	193-39-5	0.017	0.0057			
Naphthalene	91-20-3	0.176	0.059			
Phenanthrene	85-01-8	0.204	0.068			
Pyrene	129-00-0	0.195	0.065			
Metals by SW-846 6020A/7471B						
Aluminum	7429-90-5	7700	2600			
Antimony	7440-36-0	2	0.67			
Arsenic	7440-38-2	0.39	0.13			
Barium	7440-39-3	48	16			
Beryllium	7440-41-7	16	5.3			
Cadmium	7440-43-9	0.99	0.33			
Calcium	7440-70-2					
Chromium	7440-47-3	0.29	0.097			
Cobalt	7440-48-4	2.3	0.77			
Copper	7440-50-8	31.6	11			
Iron	7439-89-6	5500	1800			
Lead	7439-92-1	35.8	12			
Magnesium	7439-95-4					
Manganese	7439-96-5	180	60			
Mercury	7439-97-6	0.18	0.06			
Nickel	7440-02-0	22.7	7.6			
Potassium	7440-09-7					
Selenium	7782-49-2	2	0.67			
Silver	7440-22-4	1	0.33			
Sodium	7440-23-5					
Thallium	7440-28-0	0.078	0.026			
Vanadium	7440-62-2	39	13			
Zinc	7440-66-6	121	40			

#### **ATTACHMENT NO. 2**

#### **STATEMENT OF WORK/PRICE TABLES**

#### **REVISION 1**

## TECHNICAL SPECIFICATION FOR LABORATORY SERVICES TANK FARM 2 and 3, NAVAL STATION (NAVSTA) NEWPORT NEWPORT, RHODE ISLAND

COMPREHENSIVE LONG-TERM ENVIRONMENTAL ACTION - NAVY (CLEAN)
CONTRACT N62470-08-D-1001, CONTRACT TASK ORDER (CTO) NO. WE30 and WE59

### REMEDIAL INVESTIGATION CHEMICAL ANALYSES

This subcontract Revision 1 is issued to support a Remedial Investigation at Tank Farms 2 and 3, Naval Station (NAVSTA) Newport. Requested changes include the addition of the Tank Farm 3 analytical services to the adjoining Tank Farm 2 analytical work and revisions/additions to the original Tank Farm 2 analytical scope of work.

Tank Farm 2 changes involve the addition of PCB and lead analysis for soil and water samples and to change the extractable total petroleum hydrocarbon (ExTPH) analysis so that the petroleum hydrocarbon range is from  $C_8$ - $C_{44}$ , instead of  $C_9$ - $C_{36}$ . The Tank Farm 3 work involves the following analyses for soil: volatile organic compounds (VOCs), polycyclic aromatic hydrocarbons (PAH), PCBs, TAL metals GRO and ExTPH. The Tank Farm 3 work for aqueous samples includes VOCs, 1,2-dibromoethane (EDB), PAHs, PCBs TAL Metals, ExTPH and GRO. The Tank Farm 3 work for sediment samples includes VOCs, PAHs TAL Metals, GRO and ExTPH.

The approximate number of samples to be submitted, the type of analyses to be conducted, and the analytical methods to be used for the Remedial Investigation at Tank Farm 3, and for the changes to analytical requirements at Tank Farm 2, are summarized in Table 1.

#### All other subcontract requirements remain unchanged except as described below.

The sampling at Tank Farms 2 and 3 is scheduled for several weeks in the spring of 2013. The dates of sample collection will be communicated to the laboratory at least 2 weeks in advance.

For completeness, the project screening levels (PSLs) for the entire project (including work covered in the original scope of work) have been provided in Attachment A. Please note that some of the PSLs have changed from the original scope of work.

For the sediment non-volatile analyses, the laboratory must decant any standing water, homogenize the sample, and determine the percent moisture before sample analysis. The sample aliquots must be increased to compensate for the moisture content of the samples or the sample may be centrifuged to eliminate more water. The project screening levels listed in Attachment A must be met regardless the moisture content of the sediment samples.

If the percent moisture content is too high and the sediment samples contain noticeable organic material, further dewatering must be performed at the laboratory prior to sample analysis. This could be accomplished by freeze-drying under controlled conditions proven to recover the analysis-specific target compounds; centrifugation and decanting free water; low temperature oven drying (below 60°C); or other procedure proposed by the laboratory and approved by Tetra Tech.

## REVISION 1 - TECHNICAL SPECIFICATION FOR LABORATORY SERVICES CONTRACT N62470-08-D-1001, CTO WE30 and WE59 TANK FARM 2 and Tank Farm 3, NAVSTA NEWPORT, NEWPORT, RI REMEDIAL INVESTIGATION, CHEMICAL ANALYSES PAGE 2

Soil and sediment samples for VOC analysis will be collected using a coring device (cut off syringe). The following aliquots will be collected:

- Two 40-ml VOA vials pre-preserved with 1 g NaHSO<sub>4</sub> in 5 ml VOC-free reagent water w/ a magnetic stir bar.
- o One 40-ml VOA vial pre-preserved with 5 ml of methanol
- o One 2-oz wide-mouth jar for VOC percent moisture

The pre-preserved VOC soil and sediment sample containers must be weighed accurately to within 0.01 grams and identified with a unique ID number. Both the ID number and the applicable vial weight must be recorded on a weight tracking form for return shipment to the laboratory. When samples are received at the laboratory, the pre-preserved vials must be re-weighed and these values recorded in the weight tracking form. Tetra Tech must be contacted immediately if leaking vials are received at the laboratory.

The VOC low-concentration analysis (bisulfate-preserved vials) must be performed first for all of the soil and sediment samples in order to meet the required project screening levels (PSLs) (Attachment A-R1). If any target analyte is above the calibration range, the laboratory should perform a dilution analysis using a methanol-preserved vial, and the medium level (methanol-preserved) trip blank associated with that sample must also be analyzed.

The analytical requirements and approximate number of samples to be submitted for analysis are detailed in Table 1.

It is a requirement of Tetra Tech that the associated PDF and hard copy data packages for PCB and EDB analyses must meet Contract Laboratory Program (CLP) format, reporting, and PDF/hard copy data package deliverable requirements. Second-source initial calibration verification results must be reported on a summary form; and for the GC/MS analysis.

Attachment A-R1 details the required target analyte list and PSLs for soil, groundwater, and sediment that must be met for the Remedial Investigation at Tank Farms 2 and 3. Attachment A-R1 also presents the laboratory's soil and groundwater LOQs, LODs, and DLs as previously submitted to Tetra Tech in support of the Tank Farm 2 project. The LOQs, LODs, and DLs for soil are entered in the sediment table. The laboratory must indicate in its response whether its LOQs, LODs, and DLs for sediment are the same as for soil.

Sediment samples must be reported on a dry-weight basis.

The holding times for PCBs and EDB are as follows:

Analyses	Preservation	Holding Time
PCBs	Soil and aqueous: Cool to < 6 °C	30 days to extraction, 40 days to analysis
EDB	Aqueous: HCl to pH < 2, no headspace, cool to ≤ 6 °C	14 days to analysis

The holding times for VOCs, PAHs, and metals for sediment samples are the same as those listed in the original subcontract for soil samples.

REVISION 1 - TECHNICAL SPECIFICATION FOR LABORATORY SERVICES CONTRACT N62470-08-D-1001, CTO WE30 and WE59 TANK FARM 2 and Tank Farm 3, NAVSTA NEWPORT, NEWPORT, RI REMEDIAL INVESTIGATION, CHEMICAL ANALYSES PAGE 3

The electronic (TXT) deliverables, one PDF (CD) copy of the analytical data package, and a copy of the chain-of-custody form, should be sent to Ms. Tobrena Sedlmyer. The contact information for Ms. Sedlmyer is as follows:

Tetra Tech NUS, Inc. 661 Andersen Drive, Foster Plaza 7 Pittsburgh, PA 15220-2745

Phone: 412-921-8582 Fax: 412-921-4040

e-mail: tobrena.sedlmyer@tetratech.com

Please confirm the laboratory's ability to perform the methodologies requested at the analyte quantitation limits indicated. Also confirm available laboratory capacity, and complete/confirm the costing information indicated in Table 1-R1. All costing information must reflect the terms and conditions established by the 2010 CLEAN BOA.

#### TABLE 1 NUMBER OF SAMPLES/ANALYTICAL METHODS CTO WE59, TANK FARM 3 NAVSTA NEWPORT, NEWPORT, RI DATA GAPS ASSESSMENT, CHEMICAL ANALYSES

Matrix	Parameter <sup>(1)</sup>	Method	# Samples	Unit Price	Total Cost
	VOCs	SW-846 5035A/8260B	22	\$	\$
	PAHs	SW-846 3540C or 3550C/ 8270D SIM	18	\$	\$
	PCBs	SW-846 3540C, 3545A, or 3550C/8082A	36	\$	\$
Soil	TAL Metals	SW-846 3050B/ 6020A/7471B	36	\$	\$
	GRO	SW-846 5035A, 8015C/CA-316	36	\$	\$
	ExTPH	SW-846 3540C or 3550C, 8015C/CA-315, CA-527, CA-535, FLA-PRO	36	\$	\$
	VOCs	SW-846 5030B/8260B/ CA-202	6	\$	\$
	EDB	SW-846 8011 / CA-391	4	\$	\$
Aqueous (Groundwater	PAHs	SW-846 3510C or 3520C/ 8270D SIM/ CA-315, CA-520	5	\$	\$
and Rinsate Blanks)	PCBs	SW-846 3510C or 3520C, 8082A/ CA-329, CA-515	7	\$	\$
	TAL Metals	SW-846 3010A,6020A, 7470A/ CA-604, CA-615, CA-627	8	\$	\$
	ExTPH	SW-846 3510C or 3520C, 8015C/ CA-315, CA-520; FLA-PRO	3	\$	\$
	GRO	SW-846 5030B, 8015C / CA-316	3	\$	\$
	VOCs	SW-846 5035A/8260B	11	\$	\$
	PAHs	SW-846 3540C or 3550C/ 8270D SIM	10	\$	\$
Sediment	TAL Metals	SW-846 3050B/ 6020A/7471B	10	\$	\$
	GRO	SW-846 5035A, 8015C/CA-316	10	\$	\$
	ExTPH	SW-846 3540C or 3550C, 8015C/CA-315, CA-527, CA-535, FLA-PRO	10	\$	\$
Additional sets of pre-preserved VOC containers for 24 soil samples <sup>(2)</sup>					\$

<sup>(1)</sup> See list of required target analytes in Attachment A.(2) See text above for aliquots to be collected.

**TOTAL COST \$** 

#### NUMBER OF SAMPLES/ANALYTICAL METHODS CTO WE30, TANK FARM 2 NAVSTA NEWPORT, NEWPORT, RI REMEDIAL INVESTIGATION, CHEMICAL ANALYSES

Matrix	Parameter	Method or Modification	# Samples	Unit Price	Total Cost
Soil	PCBs	SW-846 3540C, 3545A or 3550C, 8082A/CA-329, CA-500, CA-524, CA-537	18	\$	\$
	Lead	SW-846 3050B, 6020A, 7471B/ CA-605, CA-611, CA-627	18	\$	\$
	ExTPH	Addition of the FLA-PRO method to the already contracted ExTPH analyses to get to C <sub>44</sub> .	14	\$	\$
Aqueous (Rinsate	PCBs	SW-846 3510C or 3520C, 8082A/ CA-329, CA-515	1	\$	\$
Blanks)	Lead	SW-846 3010A, 6020A, 7470A/ CA-604, CA-615, CA-627	1	\$	\$

TOTAL COST \$

Name of Laboratory_		
•		
Signature		

#### ATTACHMENT A

PROJECT-REQUIRED TARGET ANALYTES AND PROJECT SCREENING LEVELS (See tables for soil, groundwater, and sediment in accompanying Word file)

#### **ATTACHMENT NO. 2**

#### STATEMENT OF WORK/PRICE TABLES

## TECHNICAL SPECIFICATION FOR LABORATORY SERVICES TANK FARM 2, NAVAL STATION (NAVSTA) NEWPORT NEWPORT, RHODE ISLAND

### COMPREHENSIVE LONG-TERM ENVIRONMENTAL ACTION - NAVY (CLEAN) CONTRACT N62470-08-D-1001, CONTRACT TASK ORDER (CTO) NO. WE30

### REMEDIAL INVESTIGATION CHEMICAL ANALYSIS - DIOXINS

#### 1.0 INTRODUCTION

Tetra Tech NUS, Inc. (Tetra Tech) under CLEAN Contract N62470-08-D-1001, is procuring laboratory analytical services to support a Remedial Investigation at Tank Farm 2, Naval Station (NAVSTA) Newport. The requested analysis is dioxins and furans (dioxins).

The laboratory performing this analysis must be accredited by the Department of Defense Environmental Laboratory Accreditation Program (DoD ELAP) for the method and target compounds required. The laboratory must provide a copy of its ELAP accreditation letter with the bid response.

The responding laboratory must submit Limits of Quantitation (LOQs), Limits of Detection (LODs), and Detection Limits (DLs) for the analysis and matrices requested by filling out the last three columns of the tables in Attachment A and including the completed attachment with the bid response.

After award, the laboratory will be required to submit Standard Operating Procedures (SOPs) and relevant precision and accuracy limits for all preparation and analytical methods required under this scope of work. The laboratory will also be asked to complete Uniform Federal Policy (UFP) Sampling and Analysis Plan (SAP) Worksheets 19, 23, 24, 25, 26, 28, and 30 for inclusion in the SAP. The SAP will be prepared according to the UFP for Quality Assurance Project Plans (March 2005) and will utilize the 37 UFP-SAP worksheets.

#### 2.0 SAMPLE INFORMATION

The approximate number of samples to be submitted for dioxins and furans analysis and the analytical methods to be used are summarized in Table 1. This investigation includes analysis of soil samples and the associated aqueous field blank samples. The number of samples may change during the SAP review process. Significant changes in numbers of samples will be communicated to the laboratory.

The sampling is currently scheduled for several weeks in the spring or summer of 2011. The exact starting date of sample collection will be communicated to the laboratory at least 2 weeks in advance.

The samples are expected to be of low or moderate contaminant concentration. The field crew will attempt to identify any potentially high concentration samples.

Field duplicate samples will be submitted to the laboratory with "blinded" identification.

Laboratory duplicate analysis is required. The field crew will designate samples (one per twenty soil samples) for laboratory duplicate analysis. Additional volumes of these samples will be provided as necessary. The laboratory should calculate and report the Relative Percent Differences (RPDs) for the laboratory duplicate pairs.

The **2005 World Health Organization** Toxicity Equivalent Factors (TEFs) for dioxins and dioxin-like compounds must be used in Form 1DFB.

#### 3.0 ANALYSIS/REPORTING INFORMATION

One hard copy data package deliverable and two PDF CD copies must be submitted, in addition to the electronic data deliverables (EDDs) to be provided in the format described in Attachment C. The original chain-of-custody form received with the samples and signed by the laboratory sample custodian must be returned with the hard copy data package.

The analytical requirements and approximate number of samples to be submitted for analysis are detailed in Table 1. Analysis and reporting requirements addressed in the DoD Quality Systems Manual (October 2010) and the requested method must be followed. The laboratory-derived recovery limits for LCSs, and the maximum RPD for the laboratory duplicate samples must be met. Additionally, it is a requirement of Tetra Tech that the associated PDF and hard copy data packages for dioxins follow the Superfund DLM02.0 statement of work format and contain the summary forms and raw data for all field samples and laboratory quality control samples. The information in the header of the forms must be complete. The Tetra Tech sample identification numbers **must be included** on the raw data and summary forms.

Additionally, each hard copy and PDF data package must contain a summary data package. This summary data package shall consist of only the summary forms (i.e., for dioxins, Forms 1DFA through 7DFB).

Attachment A details the required target analyte list and project screening levels (PSLs), and the required LOQ Goals to be achieved by the laboratory. The laboratory must submit its LOQs, LODs, and DLs for dioxins and furans analysis for soil by filling out the last three columns of Attachment A and including the completed attachment with the bid response. If the LOQ Goal for a target analyte is not technically achievable, the laboratory should propose the lowest LOQ technically possible using the required method.

Attachment B details the required summary forms for dioxin data packages and requirements for organization/bookmarking of hard copy/PDF data packages.

The dioxins results must be reported using sample-specific Estimated Detection Limits (EDLs) per Method SW846 8290A. Non-detected results must be reported as EDL U, and results below the LOQ and above or equal to the EDL must be estimated "J". Soil sample results must be reported on a dry-weight basis.

The hard copy/PDF data package deliverable must contain a detailed case narrative for all analytical fractions. This case narrative must also include the Contract Task Order (CTO) number, the site name, and the Tetra Tech Project Manager's name. Data from all analytical runs (i.e., original, dilution, re-analysis) must be reported in the raw data and Form Is.

As part of the laboratory case narrative, it is required that the Laboratory Quality Assurance Manager sign an attestation statement verifying that all electronic diskette deliverables exactly match the data summary forms (i.e. Form Is).

As stipulated in the CLEAN Basic Ordering Agreement (BOA), Sample Delivery Group (SDG) and fractionally-specific text (TXT) files containing all environmental sample and field quality control blank analysis results must be generated in accordance with the requirements outlined in Attachment C of this specification.

Maximum holding time allowances, as defined in the following table, are to be strictly observed. Calculation of holding time is in calendar days and is to begin from the time of sample collection. The holding times are as follows:

Analyses	Preservation	Holding Time
Dioxins and Furans	Soil and Aqueous: Cool to < 6 °C	30 days to extraction; 45 days to analysis

These holding times are based on 40 CFR 136, data validation criteria, and method specific requirements, and are measured from date of collection for samples preserved as requested in the analytical methods. The holding time criteria depicted apply to all analyses necessary to successfully determine the contaminant level contained in the sample. Hence, **the holding time criteria apply to any/all subsequent sample dilutions and re-analyses**.

The Tetra Tech Project Manager for this project is Ms. Dabra Seiken, and she must be contacted in the event of any laboratory problems that could impact project deadlines (i.e., late deliverables, technical problems in the lab that could lead to late deliverables.) To insure good communication it is required that the laboratory's appointed project manager contact Ms. Seiken once a week for the entire project duration.

Contact information for Ms. Seiken is as follows:

Tetra Tech. Inc. 250 Andover Street, Suite 200 Wilmington, MA 01887 Phone: 978-474-8445

Fax: 978-474-8499

Email: dabra.seiken@tetratech.com

Technical, quality assurance, and data format concerns are to be directed to the Project Chemist, Ms. Lucy Guzman, at 978-474-8416 or via e-mail at <a href="lucy.guzman@tetratech.com">lucy.guzman@tetratech.com</a>. Ms. Guzman must be contacted and informed of any difficulties encountered during the conduct of the requested analysis.

Analytical data turnaround times are to be measured from receipt of each sample shipment. The hard copy/PDF (2 CDs) analytical data package and associated electronic (TXT) deliverables are due within the standard BOA turnaround term of 21 calendar days from receipt of the last sample in a Sample Delivery Group (SDG).

All of the samples should be reported in one SDG. The hard copy data packages, PDFs, and electronic deliverables must be received at the same time or the deliverable will be considered incomplete and payment deductions may be imposed.

The hardcopy analytical data package, one PDF (CD) copy of the analytical data package, and the original chain-of-custody form (received with the samples and signed by the laboratory sample custodian) should be sent to Ms. Lucy Guzman. The mailing address for Ms. Guzman is the same as noted above for Ms. Seiken.

The electronic (TXT) deliverables, one PDF (CD) copy of the analytical data package, and a copy of the chain-of-custody form, should be sent to Ms. Tobrena Skeen. The contact information for Ms. Skeen is as follows:

Tetra Tech, Inc. 661 Andersen Drive, Foster Plaza 7 Pittsburgh, PA 15220-2745

Phone: 412-921-8582 Fax: 412-921-4040

e-mail: tobrena.skeen@tetratech.com

#### 4.0 PERIOD OF PERFORMANCE/BOTTLEWARE INFORMATION

All samples will be shipped to the laboratory via express carrier within 48 hours of collection. The laboratory will be notified at least 2 weeks prior to sample collection.

#### Bottleware shipments will be coordinated by the field operations leader.

The laboratory is to provide all necessary sample containers (plus approximately 10% extra for breakage). All sample containers must meet ICHEM series 300 cleanliness criteria (or equivalent), and documentation of certified cleanliness must be provided. The bottleware must be shipped to the designated location in Coleman-like coolers. Each cooler must include a "temperature blank" vial. The laboratory must also provide any extra coolers needed for return shipment of samples to the laboratory for analysis. The laboratory is also requested to provide a packing slip indicating the analytical parameters for which each container type is designated, sample labels, chain-of-custody forms, and custody seals.

#### 5.0 ADDITIONAL COMMENTS/CONTACTS

The internal transfers of samples and extracts within the laboratory, must be accomplished and documented as controlled custody transfers. The laboratory must submit documentation that supports an unbroken chain of custody for samples and extracts from time of receipt or production in the laboratory until disposal.

The laboratory is to provide a minimum of 60 days storage of sample extracts and 60 intact leftover samples, as stipulated in the BOA. Additionally, the laboratory must store PDF data packages for 7 years.

All analyses conducted under this subcontract assignment are to be performed at the solicited facility only. The laboratory is not permitted to lower-tier subcontract these analyses, or analyze these samples at a corporate facility other than the facility stipulated, without prior notification and consent from the CLEAN Subcontracting Officer.

The unit cost for analysis is to include compensation for containers, preservatives, coolers, shipping costs, storage, disposal, and laboratory quality control analyses (such as laboratory duplicate and laboratory control sample analyses). **These items are not to be billed as separate line items**.

#### Contract concerns, and response to this solicitation, are to be directed to:

Ms. Meg Price CLEAN Subcontracting Officer Tetra Tech, Inc. 234 Mall Boulevard, Suite 260 King of Prussia, PA 19406

Phone: 610-491-9688 Fax: 610-491-9645

e-mail: meg.price@tetratech.com

Triplicate copies of invoices associated with the analyses contracted herein are to be submitted to the attention of the Accounting Supervisor:

Tetra Tech, Inc. 661 Andersen Drive, Foster Plaza 7 Pittsburgh, PA 15220 Phone: 412-921-8506 Fax: 412-921-4040

Please confirm the laboratory's ability to perform the methodologies requested at the analyte detection limits indicated. Also confirm available laboratory capacity, and complete/confirm the costing information indicated in Table 1. All costing information must reflect the terms and conditions established by the 2010 CLEAN BOA.

## TABLE 1 NUMBER OF SAMPLES/ANALYTICAL METHODS CTO WE30, TANK FARM 2, NAVSTA NEWPORT, NEWPORT, RHODE ISLAND REMEDIAL INVESTIGATION, CHEMICAL ANALYSIS - DIOXINS

Matrix	Parameter	Method	# Samples	Unit Price	Total Cost
Soil	Dioxins/Furans	SW-846 Method 8290A	88	\$	\$
Aqueous Field Blanks	Dioxins/Furans	SW-846 Method 8290A	4	\$	\$

TOTAL \$

Name of Laboratory_	 	
Signature		

ATTACHMENT A
PROJECT-REQUIRED TARGET ANALYTES AND PROJECT SCREENING LEVELS

## PROJECT-REQUIRED TARGET ANALYTES AND PROJECT SCREENING LEVELS CTO WE30, TANK FARM 2 NAVSTA NEWPORT, NEWPORT, RHODE ISLAND REMEDIAL INVESTIGATION, CHEMICAL ANALYSIS - DIOXINS

Matrix: Soil

		Category 1 Soil PSL <sup>(1)</sup> (mg/kg)	LOQ Goal (mg/kg)	Laboratory-Specific Limits (2)		
Analyte	CAS Number			LOQ (mg/kg)	LOD (mg/kg)	DL (mg/kg)
Dioxins and Furans by SW-8	346 8290A					
1,2,3,4,6,7,8,9-OCDD	3268-87-9	0.015	0.005			
1,2,3,4,6,7,8,9-OCDF	39001-02-0	0.012	0.004			
1,2,3,4,6,7,8-HPCDD	35822-46-9	0.00045	0.00015			
1,2,3,4,6,7,8-HPCDF	67562-39-4	0.00037	0.00012			
1,2,3,4,7,8,9-HPCDF	55673-89-7	0.00037	0.00012			
1,2,3,4,7,8-HXCDD	39227-28-6	0.000045	0.000015			
1,2,3,4,7,8-HXCDF	70648-26-9	0.000037	0.000012			
1,2,3,6,7,8-HXCDD	57653-85-7	0.000045	0.000015			
1,2,3,6,7,8-HXCDF	57117-44-9	0.000037	0.000012			
1,2,3,7,8,9-HXCDD	19408-74-3	0.000045	0.000015			
1,2,3,7,8,9-HXCDF	72918-21-9	0.000037	0.000012			
1,2,3,7,8-PECDD	40321-76-4	0.0000045	0.0000015			
1,2,3,7,8-PECDF	57117-41-6	0.00012	0.00004			
2,3,4,6,7,8-HXCDF	60851-34-5	0.000037	0.000012			
2,3,4,7,8-PECDF	57117-31-4	0.000012	0.000004			
2,3,7,8-TCDD	1746-01-6	0.0000042	0.0000014			
2,3,7,8-TCDF	51207-31-9	0.000037	0.000012			
TOTAL HPCDD	37871-00-4					
TOTAL HPCDF	38998-75-3					
TOTAL HXCDD	34465-46-8					
TOTAL HXCDF	55684-94-1					
TOTAL PECDD	36088-22-9					
TOTAL PECDF	30402-15-4					
TOTAL TCDD	41903-57-5					
TOTAL TCDF	55722-27-5					

#### Notes:

<sup>(1)</sup> PSL = Project Screening Level. PSLs are still under development and may be different in the final SAP.

<sup>(2)</sup> Laboratory to enter laboratory-specific limits. LODs and DLs will be presented in the SAP for informational purposes. Laboratory will report results using sample-specific Estimated Detection Limits (EDLs) per Method SW846 8290A. Non-detected results must be reported as EDL U, and results below the LOQ and above or equal to the EDL must be estimated "J".

<sup>-- =</sup> Not available or not applicable.

ATTACHMENT B
DATA PACKAGE DELIVERABLES REQUIREMENTS

#### DATA PACKAGE DELIVERABLE REQUIREMENTS

The laboratory is to provide two compact disks (CDs) containing a PDF file in the following format:

- 1. Table of Contents
- 2. Case Narrative
- 3. Chain-of-Custody
- 4. Data Summary Package (contains summary of all CLP or CLP like Forms 1DFA through 7 DFB per analytical fraction)
- 5. Analytical Fractions (dioxins)
  - a. Results and QC Summary (Forms 1DFA through 7 DFB for dioxins)
  - b. Raw Sample Data (includes all sample dilutions, sample re-analyses, QC samples, etc.)
  - c. Calibration Data (includes all initial and continuing calibrations)
  - d. Miscellaneous (includes extraction forms, IDLs, MDLs, etc.)

Each of the above sections should be bookmarked in the PDF for easy access.

#### Summary Form Requirements for PDF dioxin data package deliverable for non-CLP Methods:

The data package deliverable forms required for dioxins by method SW-846 8290A include:

1DFA - FORM I-HR- CDD-1 1DFB – FORM I-HR CDD-2 1DFC – FORM I-HR CDD-3 2DF – FORM II HR CDD	One Sample per summary page. Results for both method blanks and environmental samples, date of collection, preparation, and analysis. Environmental samples should be identified with the field identification numbers on the COCs. Second column confirmation and total homologue concentration summary.
<b>Lab Control Summary Form:</b> 3DFA – FORM III HR CDD	Report results for LCS analysis
Method Blank Summary Form: 4DFA – FORM IV HR CDD	Report results for method blank analysis
5DFA – FORM V HR CDD-1	Report results for window defining mix that precedes the calibration standards on each GC column and instrument use for analysis
	Report results for chromatographic resolution standard mixes on the specific GC column used
FORC FORM VIII COD 3	Report results the sequence of analyses including the WDM, isomer specificity check, calibration standards, blanks, samples, and LCS
Factor Summary:	Report the calibration response factors for each target analyte, labeled compound, and clean up standard calculated from the initial calibration
Initial Calibration Ion Abundance Ratio Summary: 6DFB – FORM VI HR CDD-2	Report the ion abundance for each of the calibration standards
Continuing Calibration Summary: 7DFA – FORM VII HR CDD-1	Report results of the 12-hour continuing calibration standard

Continuing Calibration Retention Time Summary: 7DFB – FORM VII HR CDD-2

Report retention time results of the 12-hour continuing calibration standard

### ATTACHMENT C ELECTRONIC DATA DELIVERABLE REQUIREMENTS

#### **ELECTRONIC DATA FORMAT REQUIREMENTS**

#### 1.0 INTRODUCTION

The laboratory is to submit text-based tab delimited EDD files for each SDG using Tetra Tech's laboratory data checker explained below. The files must be in the format specified in this Attachment. Additional information such as laboratory name, project name, fractions included, project number, site name/number, laboratory contact person and any specific comments related to the EDD should be included in the comments section of the EDD Submittal page.

The RESULT for nondetects should be populated with the project-specific sample quantitation reporting limits (i.e., either the sample quantitation limit or method detection limit, as specified in Section 3.0 of this scope of work. Any corrections made to the hardcopy data must also be made to the electronic file. Appropriate qualifiers as identified by the analytical protocol must also be designated; laboratory QC non-compliance codes are not to be depicted.

Tetra Tech's electronic EDD format follows the ADAPT structure and requires the A1 and A3 files. The A2 file is only required if the project is using ADAPT; and, for non-ADAPT EDD submittals the A2 file may be omitted. The EDD consists of separate, tab-delimited ASCII text files. Each file corresponds to a database table. The tables are identified as the Analytical Results Table (A1) and Sample Analysis Table (A3). A separate set of text files must be created and submitted for each sample delivery group (SDG). The files must be identified to correspond to the (A1) table and the (A3) table. The file naming convention is: the Sample Delivery Group (SDG) followed by the table identifier (A1 or A3), followed by the ".txt" extension. The file names must not contain spaces or special characters. For example, the EDD file names for a laboratory-reporting batch identified as SDG001 would be as follows:

SDG001A1.txt SDG001A3.txt

On certain projects Tetra Tech will utilize the ADAPT Electronic Data Validation software, which will require the laboratory to use the ADAPT electronic data deliverable checker software prior to submitting the files through Tetra Tech's laboratory data checker (this will be clearly specified in the Tetra Tech laboratory statement of work). The ADAPT checker software can be downloaded from Laboratory Data Consultants' web site: http://www.lab-data.com. For projects which Tetra Tech is using the ADAPT software, Tetra Tech will provide the laboratory with the project library. The laboratory is not permitted to modify the project library. ADAPT projects will require the laboratory to export all three checked files (A1, A2, and A3) from the ADAPT software and submit them through Tetra Tech's laboratory data checker. ADAPT error logs generated must be included with the electronic PDF data validation package and cannot be submitted through the laboratory data checker.

The values reported in the EDD text files must agree exactly with the final values reported on the PDF data package sample result summaries. The details of file naming conventions, data structure and data checker use are discussed below.

#### **Analytical Results Table (A1 File)**

The Analytical Results table contains analytical results and related information for target analytes in field samples and associated laboratory quality control samples (excluding calibrations and tunes). Field samples and laboratory method blanks must report a result record for each analyte reported within a method. Laboratory control samples (LCS and LCSD) and matrix spike samples (MS and MSD) must report a result record for every analyte specified as a spiked analyte in the laboratory statement of work. Table A1 in this document lists the field names and data type descriptions for the Analytical Results Table (A1).

#### Lab Instrument Table (A2 File)

A2 file is only required if the project is using ADAPT. In all other EDD submittals, the A2 file may be omitted. Laboratories should refer to the ADAPT User Guide for populating the A2 Table.

#### Sample Analysis Table (A3 File)

The Sample Analysis table contains information specific to field environmental samples and laboratory quality control analyses (excluding calibrations and tunes). A sample record must exist for each sample/method/matrix/analysis type combination. Table A3 in this document lists the field names and data type descriptions for the Sample Analysis Table (A3).

All electronic data deliverables are due within the same time established for the associated hardcopy data packages.

In addition, the laboratory QC officer must read and sign a copy of the Quality Assurance Review Form displayed on the next page of this Attachment. Electronic deliverables are not considered to be complete without the accompanying Quality Assurance Review Form.

all electronic deliverable hardcopy data. The end figures), completeness a	, as the designate in the designate in the designate in the laboratory which deliverables that have be	iewed and are in agre been reviewed for acc vill be responsible for a en found to be in error	eement with the associated curacy (including significant any labor time necessary to r. I can be reached at
( <u>)</u> deliverables.	if there are any quest	tions or problems wit	th the enclosed electronic
deliverables.			
Signature:	Title:		Date:

#### 2.0 EDD Field Properties

Tables A1 and A3 in this document specify the EDD field properties. Laboratories should refer to the ADAPT User Guide for populating the A2 Table. These include the field name, sequence order, field description, data type/length and reporting requirement for each field. Fields in the EDD **must** be sequenced according to the order that they appear below in Tables A1 and A3. For example, in the Analytical Results table (A1), the field "ClientSampleID" will always be the first piece of information to start every new line of data (or database record), followed by the field "LabAnalysisRefMethodID", "AnalysisType", etc.

When creating an EDD as a text file, use the ASCII character set in a file of lines terminated by a carriage return and line feed. No extra characters are allowed at the end of a line, after the carriage return and line feed. Enclose each data value with double quotes (text qualifier) and separate each field value with a **tab** character (tab delimiter). Data fields with no information (null) may be represented by two consecutive tabs. For example, in the Sample Analysis table, since the "Collected", "ShippingBatchID", and "Temperature" fields do not apply to laboratory generated QA/QC samples, the record for a Laboratory Control Sample by Method 8270C would be entered as follows. Note that the first two fields ("ProjectNumber" and "ProjectName") are omitted in this example.

..."LCSW100598" ` "AQ" "LCSW100598" "LCS" "8270C",...etc.

If a field is populated with less than the maximum allowed number of characters, do not pad the values with leading or trailing spaces. In the above example, although the "MatrixID" field can accommodate up to 10 characters, only 2 characters were entered in this field. **Do not include the delimiter (tab character) within any of the field values.** Example EDD files may be downloaded from the LEDD Checker application.

An example database shall be sent for review prior to the first electronic deliverable in the required .txt format. The example file will be examined for completeness and comments will be sent to the laboratory. Any questions regarding the electronic deliverable should be directed to LabSupport@tetratech.com

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

Contains laboratory test results and related information for field and QC samples (excluding instrument calibrations) on an analyte level for environmental chemistry including radiochemistry

Field Name	Field Name Description	Field Type	Field Length	Required Value
ClientSampleID	Client or contractor's identifier for a field sample	Text	25	X
	If a sample is analyzed as a laboratory duplicate, matrix spike, or matrix spike duplicate, append suffixes DUP, MS and MSD respectively to the Client Sample ID			

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiochemistry				
Field Name	Field Name Description	Field Type	Field Length	Required Value
	with no intervening spaces or hyphens (i.e. MW01DUP, MW01MS, and MW01MSD). For Method Blanks, LCS, and LCSD enter the unique LaboratorySampleID into this field.			
	Do not append suffixes to the ClientSampleID for dilutions, reanalyses, or re-extracts (the AnalysisType field is used for this distinction). For example, MW01 <u>DL</u> and MW01 <u>RE</u> are not allowed.			
	Parent sample records must exist for each MS and MSD. If an MS/MSD is shared between two EDDs, records for the MS/MSD and its parent sample must exist in the Analytical Results table for both EDDs.			
LabAnalysisRefMetho dID	Laboratory reference method ID. The method ID may be an EPA Method number or a Lab Identifier for a method such as a SOP Number, however; method ID is specified by the project. The method ID must be entered into the standard list.	Text	25	X
AnalysisType	Defines the analysis type (i.e., Dilution, Reanalysis, etc.). This field provides distinction for sample result records when multiple analyses are submitted for the same sample, method, and matrix; for example dilutions, re-analyses, and re-extracts.	Text	10	X
LabSampleID	Laboratory tracking number for field samples and lab generated QC samples such as method blank, LCS, and LCSD. There are no restrictions for the	Text	25	X

Table A1 Field Descriptions for the Analytical Results Table (Table A1)

radiochemistry					
Field Name	Field Name Description	Field Type	Field Length	Required Value	
	LabSampleID except for field length and that the LabSampleID must be distinct for a given field sample or lab QC sample and method.				
	Suffixes may be applied to the LabSampleID to designate dilutions, reanalysis, etc.				
LabID	Identification of the laboratory performing the analyses.	Text	7	X	
ClientAnalyteID	CAS Number or unique client identifier for an analyte or isotope.  If a CAS Number is not available, use a unique identifier provided by the client or contractor. The ClientAnalyteID for a particular target analyte or isotope should be specified by the project and must exist in the standard value tables for Analytes.  For the LCS, LCSD, MS, and MSD, it is only necessary to report the compounds designated as spikes in the library (and surrogates for organic methods.)  For TICs from GC/MS analyses, enter the retention time in decimal minutes as the Client Analyte ID.	Text	12	X	
AnalyteName	Chemical name for the analyte or isotope. The project specifies how an analyte or isotope is named. The analyte name must be associated to a ClientAnalyteID in the	Text	60	X	

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiochemistry			<b>-</b> :	ъ
Field Name	Field Name Description	Field	Field Length	Required Value
Field Name	standard values table for Analytes (excluding compounds designated as TIC's).	Type	Length	value
Result	Entries must be numeric. For non-detects of target analytes or isotopes and spikes, do not enter "ND" or "0". Do not leave this field blank. If an analyte or spike was not detected, enter the associated value specified in Section 3.0 of this scope of work (e.g., LOD, SQL, PQL, etc.), corrected for dilution and percent moisture as applicable. Do not enter "0". A "0" result may be acceptable for surrogate or internal standard percent recoveries; however, it should not be reported for any target compound.	Numeric (1)	20(6)	X
ResultUnits	The units defining how the values in the Result, DetectionLimit, and ReportingLimit fields are expressed. For radiochemistry this also includes how the value in the Error field is expressed.	Text	10	X
LabQualifiers	A string of single letter result qualifiers assigned by the lab based on client-defined rules and values.  The "U" Lab Qualifier must be entered for all non-detects. Other pertinent lab qualifiers may be entered with the "U" qualifier. Order is insignificant. Lab qualifiers other than those listed in the standard values table	Text	7	Q

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiocnemistry		Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
	may be used. If so, these must be added to the standard value table in the application.			
DetectionLimit	For radiochemistry methods, the minimum detectable activity for the isotope being measured.	Numeric (1)	10(6)	X
	For all other methods: The minimum detection limit value for the analyte being measured.			
	For surrogates, internal standards, etc. where detection limits are not applicable use the value -99.			
DetectionLimitType	Specifies the type of detection limit (i.e., MDA, MDL, IDL, etc.).	Text	10	X
	If -99 is specified in the DetectionLimit field us the value NA.			
RetentionTime or Error	For radiochemistry methods only, enter the 2 Sigma Counting Errors. The units for error are entered in the ResultUnits field.	Text	5	T
	For GC/MS methods only, enter the time expressed in decimal minutes between injection and detection for GC/MS TICs only			
	For target analytes in all other methods, leave this field blank. Note: GC retention times are not evaluated at this time.			
AnalyteType	Defines the type of result, such as tracer, surrogate, spike, or target compound.	Text	7	X
PercentRecovery	For radiochemistry methods: The tracer yield, if applicable.	Numeric <sup>(1)</sup>	5(3)	X

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiochemistry					
Field Name	Field Name Description	Field Type	Field Length	Required Value	
	For all other analytical methods: The percent recovery value of a spiked compound or surrogate.				
	If the spike or surrogate was not recovered because of dilution, enter "DIL". If a spike or surrogate was not recovered because of matrix interference, enter "INT". If a spike or surrogate was not recovered because it was not added to the sample, enter "NS".				
RelativePercentDiffere nce	The relative percent difference (RPD) of two QC results, such as MS/MSD, LCS/LCSD, and Laboratory Duplicates. Report RPD in Laboratory Duplicate, LCSD, and MSD records only.	Numeric <sup>(1)</sup>	5(3)	X	
ReportingLimit	Reporting limit value for the measured analyte or isotope Factor in the dilution factor and percent moisture correction, if applicable. The Reporting Limit for each analyte and matrix in a given method is specified in the project library or QAPP.  For surrogates, internal standards, etc. where reporting limits are not applicable use the value -99.	Numeric <sup>(1)</sup>	10(6)	X	
ReportingLimitType	Specifies the type of reporting limit (i.e., CRQL, PQL, SQL, RDL, etc). The Reporting Limit Type for each method and matrix is specified in the project library or QAPP.  If -99 is specified in the ReportingLimit field us the value NA.	Text	10	X	

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiochemistry		Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
ReportableResult	This field indicates whether or not the laboratory chooses an individual analyte or isotope result as reportable. Enter "YES" if the result is reportable. Enter "NO" if the result is not reportable.	Text	3	X
	If only one analysis is submitted for a particular sample and method, enter "YES" for all target compounds (where Analyte Type = TRG). For GC/MS methods enter yes for tentatively identified compounds (where Analyte Type = TIC).			
	If two or more analyses are submitted for a particular sample and method (i.e. initial analysis, reanalysis and/or dilutions), enter "YES" from only one of the analyses for each target compound. For example: a sample was run a second time at dilution because benzene exceeded the calibration range in the initial, undiluted analysis. All target analytes are reported in each analysis. For the initial analysis, (Analysis Type = RES), enter "NO" for benzene and enter "YES" for all other compounds. For the diluted analysis (Analysis Type = DL), enter "YES" for benzene and enter "NO" for all other compounds.			
	For TICs (Analyte Type = TIC), if more than one analysis is submitted for a particular sample and method, choose only one of the analyses where Reportable Result = YES for <u>all</u> TICs. For example, a sample was run a second time because one or more target compounds exceeded the calibration			

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiochemistry		Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
	range in the undiluted analysis. Choose a particular analysis and enter "YES" for all TICS. In the other analysis enter "NO" for all TICs.			
	Note that it is not necessary to report the full target analyte list for the initial result, dilution, re-analysis, or re-extraction. However, each target analyte must be reported YES once and once only in the case of multiple analyses for a given sample, method, and matrix. In the case of organics, all surrogates must be reported for all analyses submitted for a given sample, method, and, matrix.			
SpkConcnAdded	The spike added. This value must be reported in the same units as the result. Where (SA) in the following equation: % Recovery = (SSA-SC)/SA x 100% where: SSA is the spiked sample concentration (amount) after spiking. SC is the sample concentration (amount) before spiking. SA is the the expected increase in sample concentration (amount) as a result of spiking. This value must incorporate all correction factors such as dilution factor and moisture content that are applied to the spiked sample when computing the spiked sample concentration or amount. Enter -99 where no spike was added.	Numeric <sup>(1)</sup>	10(6)	X
SpkParentSampleID	The sampleID of a sample (often called the original sample) that receives a spike aliquot to form a spiked sample such as a	Text	25	X

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiocnemistry		Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
	matrix spike. This is not the same as the ID of the spiked sample (such as a matrix spike) after spiking.			
	The result for SpkParentSampleID and the result (i.e., SpkConcnAdded) for the spiked sample are used to compute percent recovery of the analyte.			
SamplePrepInitial	The initial sample preparation volume in liters (L) for aqueous samples or grams (g) for solid samples.	Numeric (1)	20(6)	
SamplePrepFinal	The final sample preparation volume in liters (L) for aqueous samples or grams (g) for solid samples.	Numeric (1)	20(6)	***************************************
LimitOfDetection	The smallest amount or concentration of a substance that must be present in a sample in order to be detected at a 99% confidence level. In other words, if a sample has a true concentration at the LOD, there is a minimum probability of 99% of reporting a "detection" (a measured value ≥ DL) and a 1% chance of reporting a non-detect (a false negative).	Numeric <sup>(1)</sup>	10(6)	N
Comment	Add any comments or additional information specific to the analyte test result data record.	Text	200	

- X Required field.
- Q Only required if laboratory has qualified the result.
- T Only required for tentatively identified compounds by GC/MS.
  - (1) Field Length indicates decimal precision in parenthesis. For example, 5(2) = a total width of 5 numbers including a maximum of 2 decimal places.

Table A3

Field Description for the Sample Analysis (Table A3)

This table contains information related to analyses of field samples and laboratory QC samples (excluding calibrations and tunes) on a sample level for environmental chemical analyses including radiochemistry

Field Name	Field Name Description	Field Type	Field Length	Required Value
ProjectNumber	Project number assigned by the client.	Text	30	X
ProjectName	Project name assigned by the client.	Text	90	X
ClientSampleID	Client or contractor's identifier for a field sample  If a sample is analyzed as a laboratory duplicate, matrix spike, or matrix spike duplicate, append suffixes DUP, MS and MSD respectively to the Client Sample ID with no intervening spaces or hyphens (i.e. MW01DUP, MW01MS, and MW01MSD). For Laboratory QC samples (i.e. Method Blanks, LCS, and LCSD, etc.) enter the unique LaboratorySampleID into this field  Do not append suffixes to the ClientSampleID for dilutions, reanalyses, or re-extracts (the Analysis_Type field is used for this distinction). For example, MW01DL and MW01RE are not allowed  Parent sample records must exist for each MS and MSD. If an MS/MSD is shared between two EDDs, records for the MS/MSD and its parent sample must exist in the Sample Analysis table for both EDDs.	Text	25	X
Collected	Date and Time of sample collection. Refer to the date/time format at the end of this table.  Leave this field blank for Method Blank, LCS, and LCSD. For Collected values that are not applicable use the value of 00/00/0000 00:00.	Date/ Time	16*	X
MatrixID	Sample matrix (i.e., AQ, SO, etc.)	Text	10	X

Table A3

Field Description for the Sample Analysis (Table A3)

This table contains information related to analyses of field samples and laboratory QC samples (excluding calibrations and tunes) on a sample level for environmental chemical analyses including radiochemistry

Field Name	Field Name Description	Field Type	Field Length	Required Value
LabSampleID	Laboratory tracking number for field samples and lab generated QC samples such as method blank, LCS, and LCSD.  There are no restrictions for the LabSampleID except field length and that the LabSampleID must be unique for a given field sample or lab QC sample and method.	Text	25	X
QCType	This record identifies the type of quality control sample QC (i.e., Duplicate, LCS, Method Blank, MS, or MSD). For regular environmental samples, populate this field with "NM".	Text	10	Х
ShippingBatchID	Unique identifier assigned to a cooler or shipping container used to transport client or field samples. Links all samples to a cooler or shipping container. No value is required for method blanks, LCS, and LCSD.	Text	25	X
Temperature	Temperature (in centigrade degrees) of the sample as received.  The storage refrigerator or room temperature should be reported (in centigrade degrees) for laboratory QC samples (i.e. method blanks, laboratory control standards).  Use -99 if temperature is not available.  This field is not required for radiochemistry methods.	Numeric (1)	10(6)	X
LabAnalysisRefMeth odID	Laboratory reference method ID. The method ID may be an EPA Method number or	Text	25	X

Table A3 Field Description for the Sample Analysis (Table A3)

		Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
	laboratory identifier for a method such as a SOP number, however; values used for Laboratory Method IDs are specified by the project and must in the in standard value list for method IDs.			
PreparationType	Preparation Method Number (i.e., 3010A, 3510C, 3550C, 5030B, etc.)  For analytical procedures that do not have a specific preparation method number, use "Gen Prep".	Text	25	X
AnalysisType	Defines the type of analysis such as initial analysis, dilution, re-analysis, etc. This field provides distinction for sample records when multiple analyses are submitted for the same sample, method, and matrix, for example: dilutions, re-analyses, and re-extracts.	Text	10	X
Prepared	Refer to the date/time format at the end of this table. If no sample preparation is involved enter the analysis date and time in this field. Refer to the date/time format at the end of this table.	Date/ Time	16*	X
Analyzed	Date and time of sample analysis. Refer to the date and time format at the end of this table. For Analyzed values that are not applicable use the value of 00/00/0000 00:00.	Date/ Time	16*	X
LabID	Identification of the laboratory performing the analysis.	Text	7	X
QCLevel	The level of laboratory QC associated with the analysis reported in the EDD. If only the Analytical Results Table (A1) and the Sample	Text	6	X

Table A3 Field Description for the Sample Analysis (Table A3)

analyses including	radiochemistry	Field	Field	Doguirod
Field Name	Field Name Description	Type	Length	Required Value
	Analysis Table (A3) information are submitted for the sample, enter "COA". If the Laboratory Instrument Table (A2) information is also submitted for the sample, enter "COCAL"			
ResultBasis	Indicates whether results associated with this sample records are reported as wet or percent moisture corrected. Enter "WET" if results are not corrected for percent moisture. Enter "DRY" if percent moisture correction is applied to results. For aqueous samples, enter "WET". For other matrices where basis is not applicable enter "NA"	Text	3	X
TotalOrDissolved	This field indicates if the results related to this sample record are reported as a total or dissolved fraction. If not applicable please report "NA"	Text	3	X
Dilution	Dilution of the sample aliquot. Enter "1" for method blanks, LCS, and LCSD, or if the field samples was analyzed without dilution.	Numeric (1)	10(6)	X
HandlingType	Indicates the type of leaching procedure, if applicable (i.e., SPLP, TCLP, WET).  Enter "NA" if the sample analysis was not performed on a leachate.	Text	10	X
HandlingBatch	Unique laboratory identifier for a batch of samples prepared together in a leaching procedure (i.e., SPLP, TCLP, or WET preparation). The HandlingBatch links samples with leaching blanks.  Enter "NA" if the sample analysis was not performed on a leachate.	Text	12	X

Table A3 Field Description for the Sample Analysis (Table A3)

analyses including	ı radiochemistry	=	<b>=•</b> - 1 1	D
Field Name	Field Name Description	Field Type	Field Length	Required Value
LeachateDate	Date and time of leaching procedure (i.e., date for SPLP, TCLP, or WET preparation). Refer to the date and time format at the end of this table.  For Analyzed values that are not applicable use the value of 00/00/0000 00:00	Date /Time	16*	X
Percent_Moisture	For soil and sediment samples, enter the percent of sample composed of water. For aqueous samples enter "100". For other matrices where Percent_Moisture is not applicable use a value of -99	Numeric <sup>(1)</sup>	10(6)	X
MethodBatch	Unique laboratory identifier for a batch of samples of similar matrices analyzed by one method and treated as a group for matrix spike, matrix spike duplicate, or laboratory duplicate association  The method batch links the matrix spike and/or matrix spike duplicate or laboratory duplicates to associated samples. Note the MethodBatch association may coincide with the PreparationBatch association. The MethodBatch is specifically used to link the MS/MSD and/or DUP to associated samples.	Text	12	X
PreparationBatch	Unique laboratory identifier for a batch of samples prepared together for analysis by one method and treated as a group for method blank, LCS and LCSD association.  The PreparationBatch links method blanks and laboratory control samples (blank spikes) to associated samples. Note, the PreparationBatch association may coincide with the MethodBatch association but the	Text	12	X

Table A3 Field Description for the Sample Analysis (Table A3)

analyses including	y radiochemistry	Field	Field	Doguirod
Field Name	Field Name Description	Type	Length	Required Value
	PreparationBatch specifically links the Method Blank and LCS to associated samples.	Type		Value
RunBatch	For all other methods the RunBatch is the unique identifier for a batch of analyses performed on one instrument under the control of one initial calibration and initial calibration verification. The RunBatch links both the initial calibration and initial calibration verification to subsequently analyzed and associated continuing calibrations, field samples, and QC analyses. For GC/MS methods, the RunBatch also links a BFB or DFTPP tune. A distinct RunBatch must used with every new initial calibration within a method  The value entered in this field links a particular sample/method/analysis type	Text	12	X
	record to a set of associated initial calibration and initial calibration verification records from Table A2.  If Table A2 is not submitted enter a value of 'NA" in this field.			
AnalysisBatch	For radiochemistry methods leave this field blank.  For all other methods the AnalysisBatch is the unique identifier for a batch of analyses performed on one instrument and under the control of a continuing calibration or continuing calibration verification. The AnalysisBatch links the continuing calibration	Text	12	X

Table A3

Field Description for the Sample Analysis (Table A3)

This table contains information related to analyses of field samples and laboratory QC samples (excluding calibrations and tunes) on a sample level for environmental chemical analyses including radiochemistry

analyses including	adioonion y	Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
	or calibration verification to subsequently analyzed and associated field sample and QC analyses. For GC/MS methods, the AnalysisBatch also links the BFB or DFTPP tune. A distinct AnalysisBatch must be used with every new continuing calibration or continuing calibration verification within a method			
	The value entered in this field links a particular sample/method/analysis type record to a set of associated continuing calibration records in the Laboratory Instrument table.			
LabReportingBatch	Unique laboratory identifier for the EDD. This is equivalent to the sample delivery group, lab work number, login ID, etc. The LabReportingBatch links all records in the EDD reported as one group. The value entered in this field must be the same in all records.	Text	12	X
LabReceipt	Date and time the sample was received in the lab. A time value of 00:00 may be entered. Refer to the date/time format at the end of this table.	Date/ Time	16*	X
LabReported	Date and time hard copy reported delivered by the lab. A time value of 00:00 may be entered. Refer to the date/time format at the end of this table.	Date/ Time	16*	X
Comment	Add any comments or additional information specific to the sample analysis data record.	Text	200	

С Only required for regular samples, duplicates and MS/MSDs.

#### X Required field.

- (1) Field Length indicates decimal precision in parenthesis. For example, 5(2) = a total width of 5 numbers including a maximum of 2 decimal places.
- \* Format Date and Time as MM/DD/YYYY hh:mm; where MM = two digit month, DD = two digit day, and YYYY = four digit year, hh = hour in 24 hour format, and mm = minutes.

#### 3.0 Laboratory Data Checker

The Laboratory Data Checker is a web-based application that will review Laboratory Electronic Data Deliverables (LEDDs) for adherence to Tetra Tech's EDD format requirements. EDDs will be reviewed for elements such as missing data and/or columns of data, and compliance of the data within each column to the required data types/lengths. Once an EDD passes through the checker with no errors, it must be submitted to Tetra Tech through the LEDD Checker application.

Access to the LEDD Checker application will be provided by an initial registration/approval process. An Information Systems Group (ISG) Administrator will approve requests for access. To access the site or begin the registration process, visit the ISG web site at <a href="http://isg.ttnus.com">http://isg.ttnus.com</a> and select the "Laboratory Checker" link on the left of the home page. Registered users may access the checker immediately by logging in to the system using their credentials. New users must select the "Register" button and provide all of the requested information.

After completing all fields on the registration form, select the "Submit" button to complete the request process. Upon verification by an ISG Administrator, an email notification will be sent verifying the user ID, password and account status. Forgotten passwords may be retrieved by using the "Forgot password?" link on the login page. Note that the email address that was provided for registration or password retrieval is the user ID and must be a valid e-mail address.

The general process for submitting EDD files through the LEDD Checker involves a 3-stage process that includes an upload stage, an error checking stage and a submittal stage.

Log into the LEDD Checker by typing your login credentials and select the "Login" button. The LEDD Checker home page provides a general overview of the checker functionality and EDD file format requirements. At the bottom of the home page, example EDDs are provided that may be viewed or downloaded. To download the files, right click on the link and select "Save target as" from the menu. Each LEDD Checker page includes a navigation bar with links to return to the home page or continue the checking and submittal process. Users should **NOT** use the back or forward buttons on the browser, instead use the links provided in the application to navigate through the site.

Detailed information regarding EDD preparation, formatting requirements and text file naming conventions are provided in the Electronic Data Format Requirements Section of the Laboratory SOW.

Begin the upload stage by selecting the "Upload/Check Files" link on the home page. Follow the steps on the upload page starting with the selection of the laboratory name that corresponds to your organization. If your organization is not listed, contact <a href="mailto:LabSupport@tetratech.com">LabSupport@tetratech.com</a>, and provide a full description of your organization name, contact information and include "Laboratory Contractor ID Request" in the subject line. An ISG Administrator will respond to the request via e-mail.

Load the appropriate A1, A2, or A3 target EDD files by clicking the "Browse" button next to each data table input box. A file browser dialog will appear allowing files to be selected from a local or network drive. After the EDD files are loaded, click the "Upload" button to complete the upload stage. Note that each table may be uploaded and checked separately; however, a minimum of the A1 and A3 files are required in order to submit the EDDs.

If the file upload was successful, the checking page will immediately load. Begin the checking stage by selecting the "Check Files" button. The LEDD Checker will begin validating the EDD files for compliance. Depending on file size and network activity the validation process may take several minutes. The progress should be displayed in the information bar at the bottom of the browser window. **Do not** select the "Check Files" button again or otherwise use the browser during this process. Other applications may be used; however, note that the LEDD Checker may not sit idle for more than 30 minutes. If the time is exceeded a new session must be started in a new browser window.

Any errors will be processed and returned on the error page. The following general errors may be returned.

- Column count / table structure errors due to column header names being included, improper delimiter, extra tabs, extra or missing columns of data, spaces or other characters at the end of a row.
- Row and column value specific errors may occur for one or more reasons including: data truncation, invalid date / time format, invalid decimal precision or field width exceedance, or if a value is not in a list of valid values or expected range.

If column count / table structure errors are encountered, the LEDD Checker will return an error and stop the checking process.

The EDDs will not be processed any further until the column errors are resolved. Text fields are validated for truncation. Date / Time fields are validated for truncation and format compliance. Numeric decimal fields are validated for truncation, character type compliance and decimal precision. All required fields are validated for null values or empty text strings (i.e. spaces). The LEDD Checker will return a list of all errors in and include a reference to the row number on which the error occurred. Note that consecutive EDD files may be loaded and checked, and submitted while logged in. However, no data may be submitted until all EDD files have passed through the LEDD Checker without errors. The list of errors may be printed by selecting the "Print this Page" button from the checker error page.

If the EDD files pass with no errors, the submittal page will immediately load. To complete the submittal stage, include the following information in the comment and additional information area of the form: laboratory name, laboratory contact person, project name, project number, site name/number, fractions included and any specific comments related to the EDD. Select the "Submit Files" button to continue the submittal process.

The submittal stage is not considered complete until a unique ticket key reference is returned in the browser window. The ticket key reference must be printed for record of submission and future reference. In addition, a copy of the ticket key reference must be included in the PDF data package.

#### ATTACHMENT A - TABLE A-1 SOIL PROJECT-REQUIRED TARGET ANALYTES AND PSLs SAMPLING AND ANALYSIS PLAN

#### REMEDIAL INVESTIGATION TANK FARM 2 NAVAL STATION NEWPORT, NEWPORT, RI PAGE 1 OF 2

		Category	Category	Lower of Category 1		Laborato	ry-Specific	Limits (2)
Analyte	CAS Number	1 Soil PSL <sup>(1)</sup> (mg/kg)	2 Soil PSL <sup>(1)</sup> (mg/kg)	or Category 2 Soil PSLs <sup>(1)</sup> (mg/kg)	LOQ Goal (mg/kg)	LOQ (mg/kg)	LOD (mg/kg)	DL (mg/kg)
VOCs by SW-846 8260C								
1,2,4-Trimethylbenzene	95-63-6		10000	10000	3300			
1,2-Dibromoethane (EDB)	106-93-4		0.07	0.07	0.023			
1,3,5-Trimethylbenzene	108-67-8		10000	10000	3300			
2-Butanone (MEK)	78-93-3		10000	10000	3300			
2-Hexanone	591-78-6		10000	10000	3300			
4-Methyl-2-pentanone (MIBK)	108-10-1		10000	10000	3300			
Acetone	67-64-1		10000	10000	3300			
Benzene	71-43-2		4.3	4.3	1.4			
Bromoform	75-25-2		720	720	240			
Bromomethane	74-83-9		2900	2900	970			
Carbon disulfide	75-15-0		10000	10000	3300			
Cyclohexane	110-82-7		10000	10000	3300			
Ethylbenzene	100-41-4		62	62	21			
Isopropylbenzene	98-82-8		10000	10000	3300			
m,p-Xylenes	179601-23-1		10000	10000	3300			
Methyl acetate	79-20-9		10000	10000	3300			
Methylcyclohexane	108-87-2		10000	10000	3300			
Methyl-tert-butyl ether	1634-04-4		100	100	33			
Naphthalene	91-20-3		10000	10000	3300			
n-Butylbenzene	104-51-8		10000	10000	3300			
n-Propylbenzene	103-65-1		10000	10000	3300			
o-Xylene	95-47-6		10000	10000	3300			
p-Isopropyltoluene	99-87-6		10000	10000	3300			
sec-Butylbenzene	135-98-8		10000	10000	3300			
Styrene	100-42-5		64	64	21			
tert-Butylbenzene	98-06-6		10000	10000	3300			
Toluene	108-88-3		54	54	18			
Xylenes (total)	1330-20-7		10000	10000	3300			
PAHs by 8270D SIM	1330-20-7		10000	10000	3300			
2-Methylnaphthalene	91-57-6	15	10000	15	5			
Acenaphthene	83-32-9	29	10000	29	9.7			
Acenaphthylene	208-96-8	29	10000	29	9.7			
Anthracene	120-12-7	29	10000	29	9.7			
Benzo(a)anthracene	56-55-3	0.15	7.8	0.15	0.05			
Benzo(a)pyrene	50-33-8	0.15	0.8	0.15	0.005			
Benzo(b)fluoranthene	205-99-2	0.015	7.8	0.015	0.005			
Benzo(g,h,i)perylene	191-24-2	1.1	10000	1.1	0.05			
Benzo(k)fluoranthene	207-08-9	1.1	78	1.1	0.37	1		
Chrysene	218-01-9	1.1	780	1.1	0.37	1		
Dibenzo(a,h)anthracene						}		
( · /	53-70-3	0.015	0.8	0.015	0.005	}		
Fluoranthene	206-44-0	29	10000	29	9.7			
Fluorene	86-73-7	29	10000	29	9.7			
Indeno(1,2,3-c,d)pyrene	193-39-5	0.15	7.8	0.15	0.05	1		
Naphthalene	91-20-3	0.0094	10000	0.0094	0.0031			
Phenanthrene	85-01-8	29	10000	29	9.7	1		
Pyrene	129-00-0	1.1	10000	1.1	0.37			

#### ATTACHMENT A - TABLE A-1 SOIL PROJECT-REQUIRED TARGET ANALYTES AND PSLs SAMPLING AND ANALYSIS PLAN

#### REMEDIAL INVESTIGATION TANK FARM 2 NAVAL STATION NEWPORT, NEWPORT, RI

PAGE 2 OF 2

		Category	Category	Lower of Category 1		Laborato	ry-Specific	Limits (2)
Analyte	CAS Number	1 Soil PSL <sup>(1)</sup> (mg/kg)	2 Soil PSL <sup>(1)</sup> (mg/kg)	or Category 2 Soil PSLs <sup>(1)</sup> (mg/kg)	LOQ Goal (mg/kg)	LOQ (mg/kg)	LOD (mg/kg)	DL (mg/kg)
TAL Metals by SW-846 601	OC/6020A <sup>(3)</sup> /7471	В						
Aluminum	7429-90-5	50		50	17			
Antimony	7440-36-0	0.27		0.27	0.09			
Arsenic	7440-38-2	0.026		0.026	0.0087			
Barium	7440-39-3	330		330	110			
Beryllium	7440-41-7	16		16	5.3			
Cadmium	7440-43-9	0.36		0.36	0.12			
Calcium	7440-70-2							
Chromium	7440-47-3	0.0166		0.0166	0.0055			
Cobalt	7440-48-4	2.3		2.3	0.77			
Copper	7440-50-8	28		28	9.3			
Iron	7439-89-6	5500		5500	1800			
Lead	7439-92-1	11		11	3.7			
Magnesium	7439-95-4							
Manganese	7439-96-5	180		180	60			
Mercury	7439-97-6	0.1		0.1	0.033			
Nickel	7440-02-0	38		38	13			
Potassium	2023695							
Selenium	7782-49-2	0.63		0.63	0.21			
Silver	7440-22-4	4.2		4.2	1.4			
Sodium	7440-23-5							
Thallium	7440-28-0	1		1	0.33			
Vanadium	7440-62-2	7.8		7.8	2.6			
Zinc	7440-66-6	46		46	15			
Petroleum Hydrocarbons								
GRO (C5-C12)								
ExTPH (C9-C36)								
TPH <sup>(4)</sup>			2500	2500	830			

#### Notes:

- (1) PSLs are still under development and may be different in the final SAP.
- (2) Laboratory is to enter laboratory-specific limits. If no LOQ Goal is presented or a LOQ below the LOQ Goal is not technically achieveable by the required method, the laboratory should enter its lowest limits achievable by the required method.
- (3) The laboratory must identify which metals will be analyzed by 6020A in order to meet the LOQ Goals, if technically possible.
- (4) The laboratory will analyze for GRO and ExTPH. TPH will be the sum of GRO and ExTPH.

#### Abbreviations:

ExTPH = Extractable TPH

GRO = Gasoline Range Organics

PSL = Project Screening Level

TPH = Total Petroleum Hydrocarbons

# ATTACHMENT A - TABLE A-2 GROUNDWATER PROJECT-REQUIRED TARGET ANALYTES AND PSLS SAMPLING AND ANALYSIS PLAN REMEDIAL INVESTIGATION TANK FARM 2 NAVAL STATION NEWPORT, NEWPORT, RI

PAGE 1 OF 2

		Category 2	LOQ Goal	Laborato	ry-Specifi	c Limits <sup>(2)</sup>
Analyte	CAS Number	Groundwater PSL <sup>(1)</sup> (μg/L)	(µg/L)	LOQ (µg/L)	LOD (µg/L)	DL (µg/L)
VOCs by SW-846 8260C				•		I.
1,2,4-Trimethylbenzene	95-63-6					
1,2-Dibromoethane (EDB)	106-93-4					
1,3,5-Trimethylbenzene	108-67-8					
2-Butanone (MEK)	78-93-3					
2-Hexanone	591-78-6					
4-Methyl-2-pentanone (MIBK)	108-10-1					
Acetone	67-64-1					
Benzene	71-43-2	140	47			
Bromoform	75-25-2					
Bromomethane	74-83-9					
Carbon disulfide	75-15-0					
Cyclohexane	110-82-7					
Ethylbenzene	100-41-4	1600	530			
Isopropylbenzene	98-82-8					
m,p-Xylenes	179601-23-1					
Methyl acetate	79-20-9					
Methylcyclohexane	108-87-2					
Methyl-tert-butyl ether	1634-04-4	5000	1700			
Naphthalene	91-20-3					
n-Butylbenzene	104-51-8					
n-Propylbenzene	103-65-1					
o-Xylene	95-47-6					
p-Isopropyltoluene	99-87-6					
sec-Butylbenzene	135-98-8					
Styrene	100-42-5	2200	730			
tert-Butylbenzene	98-06-6					
Toluene	108-88-3	1700	570			
Xylenes (total)	1330-20-7					
PAHs by 8270D SIM	•			•		•
2-Methylnaphthalene	91-57-6					
Acenaphthene	83-32-9					
Acenaphthylene	208-96-8					
Anthracene	120-12-7					
Benzo(a)anthracene	56-55-3					
Benzo(a)pyrene	50-32-8					
Benzo(b)fluoranthene	205-99-2					
Benzo(g,h,i)perylene	191-24-2					
Benzo(k)fluoranthene	207-08-9					
Chrysene	218-01-9					
Dibenzo(a,h)anthracene	53-70-3					
Fluoranthene	206-44-0					

#### ATTACHMENT A - TABLE A-2 GROUNDWATER PROJECT-REQUIRED TARGET ANALYTES AND PSLs SAMPLING AND ANALYSIS PLAN

#### REMEDIAL INVESTIGATION TANK FARM 2 NAVAL STATION NEWPORT, NEWPORT, RI PAGE 2 OF 2

		Category 2	LOQ Goal	Laboratory-Specific Limits (2)		
Analyte	CAS Number Groundwater PSL <sup>(1)</sup> (µg/L)		(μg/L)	LOQ (µg/L)	LOD (µg/L)	DL (µg/L)
PAHs by 8270D SIM CONTINUED						
Fluorene	86-73-7					
Indeno(1,2,3-c,d)pyrene	193-39-5					
Naphthalene	91-20-3					
Phenanthrene	85-01-8					
Pyrene	129-00-0					
Petroleum Hydrocarbons by SW-846 8	3015D					
GRO (C5-C12)						
ExTPH (C9-C36)						
TPH <sup>(3)</sup>			-			

#### Notes:

- (1) PSLs are still under development and may be different in the final SAP.
- (2) Laboratory to enter laboratory-specific limits. If no LOQ Goal is presented, the laboratory should enter its lowest limits achievable for the required method.
- (3) The laboratory will analyze for GRO and ExTPH. TPH will be the sum of GRO and ExTPH.

#### **Abbreviations:**

ExTPH = Extractable TPH

GRO = Gasoline Range Organics

PSL = Project Screening Level

TPH = Total Petroleum Hydrocarbons

#### **ATTACHMENT NO. 2**

#### STATEMENT OF WORK/PRICE TABLES

### TECHNICAL SPECIFICATION FOR LABORATORY SERVICES TANK FARM 2, NAVAL STATION (NAVSTA) NEWPORT NEWPORT, RHODE ISLAND

COMPREHENSIVE LONG-TERM ENVIRONMENTAL ACTION - NAVY (CLEAN) CONTRACT N62470-08-D-1001, CONTRACT TASK ORDER (CTO) NO. WE30

### REMEDIAL INVESTIGATION CHEMICAL ANALYSES

#### 1.0 INTRODUCTION

Tetra Tech NUS, Inc. (TtNUS) under CLEAN Contract N62470-08-D-1001, is procuring laboratory analytical services to support a Remedial Investigation at Tank Farm 2, Naval Station (NAVSTA) Newport. Requested analyses include volatile organic compounds (VOCs), polycyclic aromatic hydrocarbons (PAHs), gasoline range organics ( $C_5$ - $C_{12}$ ) (GRO), extractable total petroleum hydrocarbons ( $C_9$ - $C_{36}$ ) (ExTPH), and target analyte list (TAL) metals.

The laboratory performing these analyses must provide a copy of its Department of Defense (DOD) Environmental Laboratory Accreditation Program (ELAP) accreditation letter; the scope of the ELAP accreditation must include all methods and all analytes requested.

The responding laboratory must submit Limits of Quantitation (LOQs), Limits of Detection (LODs), and Detection Limits (DLs) for all analyses and matrices requested by filling out the last three columns of the tables in Attachment A and including the completed attachment with the bid response.

After award, the laboratory will be required to submit Standard Operating Procedures (SOPs) and relevant precision and accuracy limits for all preparation and analytical methods required under this scope of work. The laboratory will also be asked to complete Uniform Federal Policy (UFP) Sampling and Analysis Plan (SAP) Worksheets 19, 23, 24, 25, 26, 28, and 30 for inclusion in the SAP. The SAP will be prepared according to the UFP for Quality Assurance Project Plans (March 2005) and will utilize the 37 UFP-SAP worksheets.

#### 2.0 SAMPLE INFORMATION

The approximate number of samples to be submitted, the type of analyses to be conducted, and the analytical methods to be used are summarized in Table 1. This Remedial Investigation includes analysis of groundwater and soil samples. It may also include fingerprint analysis of non-aqueous phase liquid (NAPL) samples, if NAPL is found.

The sampling is scheduled for several weeks in the spring or summer of 2011. The exact date of sample collection will be communicated to the laboratory at least 2 weeks in advance.

The samples are expected to be of low or moderate contaminant concentration. The field crew will attempt to identify any potentially high concentration samples.

If groundwater samples submitted to the laboratory contain a significant layer of free product NAPL, <u>the laboratory should contact TtNUS for instructions</u> about whether to analyze both phases as separate samples.

Field duplicate samples will be submitted to the laboratory with "blinded" identification. The field crew will designate samples (one per twenty samples of like matrices) for matrix spike/matrix spike duplicate (MS/MSD) analyses (organics) or matrix spike/laboratory duplicate analyses (metals). Additional volumes of these samples will be provided as necessary.

#### Soil Volatile Analyses

Soil samples for VOC and GRO analysis will be collected using a coring device (cut off syringe). The following aliquots will be collected:

- VOC analysis
  - Two 40-ml VOA vials pre-preserved with 1 g NaHSO<sub>4</sub> in 5 ml VOC-free reagent water w/ a magnetic stir bar.
  - $\circ$  One 40-ml VOA vial pre-preserved with 5 ml of methanol
  - o One 2-oz wide-mouth jar for VOC/GRO percent moisture
- GRO analysis One 40-ml VOA vial pre-preserved with 5 ml of methanol

The pre-preserved VOC and GRO soil sample containers must be weighed accurately to within 0.01 grams and identified with a unique ID number. Both the ID number and the applicable vial weight must be recorded on a weight tracking form for return shipment to the laboratory. When samples are received at the laboratory, the pre-preserved vials must be re-weighed and these values recorded in the weight tracking form. TtNUS must be contacted immediately if leaking vials are received at the laboratory.

The VOC low-concentration analysis (bisulfate-preserved vials) must be performed first for all of the soil samples in order to meet the required LOQ Goals (Attachment A). If any target analyte is above the calibration range, the laboratory should perform a dilution analysis using a methanol-preserved vial, and the medium level (methanol-preserved) trip blank associated with that sample must also be analyzed.

#### **GRO/ExTPH Analyses**

For soil and groundwater samples, the GRO fraction must include all petroleum hydrocarbons ranging from  $C_5$  to  $C_{12}$ , and the ExTPH fraction must include petroleum hydrocarbons from  $C_9$  to  $C_{36}$ . In addition to quantitating the GRO and ExTPH concentrations, the laboratory must identify the fingerprints that are characteristic of fuel contamination present in the samples. The gas chromatograms must depict the fingerprint patterns at greater than 25 percent of the full scale.

For NAPL samples, only the fingerprint analysis (GRO and ExTPH) is required. Approximately 1 g of NAPL product will be collected. The GRO aliquots will be preserved with 5 ml of methanol.

#### 3.0 ANALYSIS/REPORTING INFORMATION

One hard copy data package deliverable and two PDF CD copies must be submitted, in addition to the electronic data deliverables to be provided in the format described in Attachment C. The original chain-of-custody form received with the samples and signed by the laboratory sample custodian must be returned with the hard copy data package.

The analytical requirements and approximate number of samples to be submitted for analysis are detailed in Table 1. Analysis and reporting requirements addressed in the Department of Defense (DOD) Quality

Systems Manual (October 2010) and the requested methods must be followed. Additionally, it is a requirement of TtNUS that the associated PDF and hard copy data packages for VOC, PAH, and metal analyses must meet Contract Laboratory Program (CLP) format, reporting, and PDF/hard copy data package deliverable requirements. Second-source initial calibration verification results must be reported on a summary form; and for the GC/MS analysis, the analytes associated with each internal standard must be identified. The PDF and hard copy data packages for the GRO and ExTPH analyses must be in a CLP-modified format and must include the appropriate summary forms and raw data for all samples and laboratory quality control samples. The summary forms should include the method-specific quality control limits (recoveries, relative percent differences, relative standard deviations, and/or percent differences, etc.). The TtNUS sample identification numbers **must be included** on the raw data and summary forms.

Additionally, each hard copy and PDF data package must contain a summary data package. This summary data package shall consist of only the summary forms (i.e., the CLP-like equivalent of Forms 1 through 15).

Attachment A details the required target analyte list and project screening levels (PSLs), and the LOQ Goals that must be met. The laboratory must submit its LOQs, LODs, and DLs for all analyses and matrices requested by filling out the last three columns of the tables in Attachment A and including the completed attachment with the bid response. If the LOQ Goal for a target analyte is not technically achievable using the required method, or if no LOQ Goal is listed for a target analyte in Attachment A, the laboratory should propose the lowest LOQ technically possible for the required method.

Attachment B details the required summary forms for CLP-like data packages and requirements for organization/bookmarking of hard copy/PDF data packages.

**Non-detected organic results and metals results** must be reported down to the laboratory's LODs. Positive results above the DL but below the laboratory's LOQ must be reported as estimated values qualified with a "J" for both organic and metals analyses. Soil samples must be reported on a dry-weight basis.

The hard copy/PDF data package deliverable must contain a detailed case narrative for all analytical fractions. This case narrative must also include the Contract Task Order (CTO) number, the site name, and the TtNUS Project Manager's name. Data from all analytical runs (i.e., original, dilution, re-analysis) must be reported in the raw data and Form Is for organic analyses. For metals analyses, only the final sample results should be reported in the Form Is, and data from all analytical runs must be included in the raw data.

As part of the laboratory case narrative, it is required that the Laboratory Quality Assurance Manager sign an attestation statement verifying that all electronic diskette deliverables exactly match the data summary forms (i.e. Form Is).

As stipulated in the CLEAN Basic Ordering Agreement (BOA), Sample Delivery Group (SDG) and fractionally-specific text (TXT) files containing all environmental sample and field quality control blank analysis results must be generated in accordance with the requirements outlined in Attachment C of this specification.

Maximum holding time allowances, as defined in the following table, are to be strictly observed. Calculation of holding time is in calendar days and is to begin from the time of sample collection. The holding times are as follows:

ΑF

Analyses	Preservation	Holding Time
	Aqueous: HCl to pH < 2, cool to < 6 °C	
VOCs	Soil (low): 1 g NaHSO₄ in 5 ml VOC-free reagent water, cool to < 6 °C	14 days to analysis
	Soil (medium): 5 ml methanol, cool to < 6 °C	
Aqueous: HCl to pH < 2, cool to < 6 °C		14 dove to englisis
GRO*	Soil/NAPL: 5 ml methanol, cool to < 6 °C	14 days to analysis
PAHs, ExTPH*	Aqueous: Cool to < 6 °C	7 days to extraction, 40 days to analysis
PARS, EXTPR	Soil/NAPL: Cool to < 6 °C	14 days to extraction, 40 days to analysis
TAL Metals	Aqueous: HNO <sub>3</sub> to pH < 2, cool to < 6 °C	6 months to analysis except for
TAL IVICIDIS	Soil: Cool to < 6 °C	mercury; 28 days to analysis for mercury

<sup>\*</sup>GRO and ExTPH analysis in soil and aqueous samples includes both quantitation and fingerprint analysis. GRO and ExTPH analysis in NAPL, if found, includes only fingerprint analysis.

These holding times are based on 40 CFR 136, data validation criteria, and method specific requirements, and are measured from date of collection for samples preserved as requested in the analytical methods. The holding time criteria depicted apply to all analyses necessary to successfully determine the contaminant level contained in the sample. Hence, **the holding time criteria apply to any/all subsequent sample dilutions and re-analyses**.

The TtNUS Project Manager for this project is Ms. Dabra Seiken, and she must be contacted in the event of any laboratory problems that could impact project deadlines (i.e., late deliverables, technical problems in the lab that could lead to late deliverables.) To ensure good communication, it is required that the laboratory's appointed project manager contact Ms. Seiken once a week for the entire project duration.

Contact information for Ms. Seiken is as follows:

Tetra Tech NUS 250 Andover Street, Suite 200 Wilmington, MA 01887 Phone: 978-474-8445

Fax: 978-474-8499

Email: dabra.seiken@tetratech.com

Technical, quality assurance, and data format concerns are to be directed to the Project Chemist, Ms. Lucy Guzman, at 978-474-8416 or via e-mail at <a href="https://lucy.guzman@tetratech.com">lucy.guzman@tetratech.com</a>. Ms. Guzman must be contacted and informed of any difficulties encountered during the conduct of the requested analyses.

Analytical data turnaround times are to be measured from receipt of each sample shipment. All hard copy/PDF (2 CDs) analytical data packages and associated electronic (TXT) deliverables are due within the standard BOA turnaround term of 21 calendar days from receipt of the last sample in a Sample Delivery Group (SDG).

The SDGs must contain no more than 20 samples. The frequency with which SDGs contain fewer than 20 samples should be minimal. The hard copy data packages, PDF files (CDs), and electronic deliverables must be received at the same time or the deliverable will be considered incomplete and payment deductions may be imposed.

The hardcopy analytical data package, one PDF (CD) copy of the analytical data package, and the original chain-of-custody form (received with the samples and signed by the laboratory sample custodian) should be sent to Ms. Lucy Guzman. The contact information for Ms. Guzman is the same as noted above for Ms. Seiken except that her direct phone number is (978) 474-8416 and her email address is lucy.guzman@tetratech.com.

The electronic (TXT) deliverables, one PDF (CD) copy of the analytical data package, and a copy of the chain-of-custody form, should be sent to Ms. Tobrena Skeen. The contact information for Ms. Skeen is as follows:

Tetra Tech NUS, Inc. 661 Andersen Drive, Foster Plaza 7 Pittsburgh, PA 15220-2745 Phone: 412-921-8582

Fax: 412-921-4040

e-mail: tobrena.skeen@tetratech.com

#### 4.0 PERIOD OF PERFORMANCE/BOTTLEWARE INFORMATION

All samples will be shipped to the laboratory via express carrier within 48 hours of collection. **The laboratory must be capable of receiving samples on Saturdays.** Please circle the Yes or No at the bottom of Table 1 which will indicate if the laboratory will provide courier service at no extra charge. The laboratory will be notified at least 2 weeks prior to sample collection.

Bottleware shipments will be coordinated by the field operation leader.

The laboratory is to provide all necessary sample containers (plus approximately 10% extra for breakage). All sample containers must meet ICHEM series 300 cleanliness criteria (or equivalent), and documentation of certified cleanliness must be provided. All of the appropriate sample bottleware must be pre-preserved. The bottleware must be shipped to the designated location in Coleman-like coolers. Each cooler must include a "temperature blank" vial. The laboratory must also provide any extra coolers needed for return shipment of samples to the laboratory for analysis. The laboratory is also requested to provide a packing slip indicating the analytical parameters for which each container type is designated, sample labels, and chain-of-custody forms.

For this project, TtNUS plans to analyze 12 soil samples for VOCs and GRO; however, the laboratory is requested to provide 78 additional sets of pre-preserved sample containers for soil interval samples

to be collected but not analyzed. The laboratory should include the cost of these additional pre-preserved sample containers as a separate line item in the price sheet (Table 1).

The laboratory must provide Material Safety Data Sheets (MSDSs) for all preservatives sent with each bottleware shipment to the field. MSDSs must be representative of the chemicals provided as preservatives with regard to mixtures and/or purity of the chemicals. For example if a 35% sulfuric acid solution is the preservative, the MSDS provided should be for 35% sulfuric acid solution not 96% sulfuric acid.

#### 5.0 ADDITIONAL COMMENTS/CONTACTS

Within the laboratory, the internal transfer of samples, extracts, and digestates must be accomplished and documented as controlled custody transfers. The laboratory must submit the documentation that supports an unbroken chain of custody for samples, digestates and extracts from time of receipt or production in the laboratory until disposal.

The laboratory is to provide a minimum of 60 days storage of sample extracts/digestates and 60 days storage of intact leftover sample aliquots, as stipulated in the BOA. Additionally, the laboratory must store PDF data packages for 7 years.

All analyses conducted under this subcontract assignment are to be performed at the solicited facility only. The laboratory is not permitted to lower-tier subcontract these analyses, or analyze these samples at a corporate facility other than the facility stipulated without prior notification and consent from the CLEAN Subcontracting Officer.

The unit cost for analysis is to include compensation for containers, preservatives, coolers, shipping costs, storage, disposal, and laboratory quality control analyses (such as matrix spike, matrix spike duplicate, laboratory duplicate, and laboratory control sample analyses.) These items are not to be billed as separate line items.

Contract concerns, and response to this solicitation, are to be directed to:

Ms. Meg Price **CLEAN Subcontracting Officer** Tetra Tech NUS. Inc. 234 Mall Boulevard, Suite 260 King of Prussia, PA 19406-1433

Phone: 610-491-9688 Fax: 610-491-9645

e-mail: meg.price@tetratech.com

Triplicate copies of invoices associated with the analyses contracted herein are to be submitted to the attention of the Accounting Supervisor:

> Tetra Tech NUS, Inc. 661 Andersen Drive, Foster Plaza 7 Pittsburgh, PA 15220

Phone: 412-921-8506

ΑF

Fax: 412-921-4040

Please confirm the laboratory's ability to perform the methodologies requested at the analyte quantitation limits indicated. Also confirm available laboratory capacity, and complete/confirm the costing information indicated in Table 1. All costing information must reflect the terms and conditions established by the 2010 CLEAN BOA.

### TABLE 1 NUMBER OF SAMPLES/ANALYTICAL METHODS CTO WE30, TANK FARM 2 NAVSTA NEWPORT, NEWPORT, RI REMEDIAL INVESTIGATION, CHEMICAL ANALYSES

Matrix	Parameter <sup>(1)</sup>	Method <sup>(2)</sup>	# Samples	Unit Price	Total Cost	
	VOCs	SW-846 5035/8260C	17	\$	\$	
	GRO (C5-C12) (3)	SW-846 5035/ 8015D	17	\$	\$	
Soil	ExTPH (C9-C36) (3)	SW-846 3540C or 3550C/ 8015D	14	\$	\$	
	PAHs	SW-846 3540C or 3550C/ 8270D SIM	146	\$	\$	
	TAL Metals	SW-846 3050B/ 6010C/6020A <sup>(4)</sup> /7471B	132	\$	\$	
	VOCs	SW-846 5030B/8260C	18	\$	\$	
Aqueous	GRO (C5-C12) (3)	SW-846 5030B/8015D	18	\$	\$	
(Groundwater and Rinsate	ExTPH (C9-C36) (3)	SW-846 3510C or 3520C/ 8015D	16	\$	\$	
Blanks)	PAHs	SW-846 3510C or 3520C/ 8270D SIM	22	\$	\$	
	TAL Metals	SW-846 3010A/ 6010C/6020A <sup>(5)</sup> /7470A	6	\$	\$	
	GRO Fingerprint	SW-846 8015D	3	\$	\$	
NAPL	ExTPH Fingerprint	SW-846 8015D	3	\$	\$	
Additional	\$	\$				
Additional	Additional sets of pre-preserved GRO containers for 78 soil samples <sup>(6)</sup>					

(1) See list of required target analytes in Attachment A. (2) Laboratory may use a different version of the SW-846 methods listed; if so, the laboratory must identify on this page and on Attachment A the method version to be used. (3) Soil and groundwater GRO and ExTPH analyses include quantitation and fingerprint analysis. (4) Laboratory must identify on Attachment A which metals will be analyzed by method 6020A in order to meet the LOQ Goals. (5) Aqueous metals analysis is for soil-associated rinsate blanks only. The rinsate blanks should be analyzed for a particular metal by the same method (6010C or 6020A) as the method to be used for soils. (6) See Section 2.0 for aliquots to be collected.

#### **TOTAL COST \$**

The laboratory must point out if it is not DOD ELAP accredited for any of the methods and/or analytes requested. Clearly state the methods and/or analytes that you are NOT accredited for (if any).

## TABLE 1 (Continued) NUMBER OF SAMPLES/ANALYTICAL METHODS CTO WE30, TANK FARM 2 NAVSTA NEWPORT, NEWPORT, RI REMEDIAL INVESTIGATION, CHEMICAL ANALYSES

Can the laboratory provide sample pick-up on site?	YES or NO (circle one
If yes is there a charge and what is that charge?	
Name of Laboratory	
Signature	

#### **ATTACHMENT A**

PROJECT-REQUIRED TARGET ANALYTES AND PROJECT SCREENING LEVELS (See tables for soil and groundwater in accompanying Excel files)

ATTACHMENT B
DATA PACKAGE DELIVERABLES REQUIREMENTS

#### DATA PACKAGE DELIVERABLE REQUIREMENTS

The laboratory is to provide a hard copy plus two compact disks (CDs) each containing a PDF file in the following format:

- 1. Table of Contents
- 2. Case Narrative
- 3. Chain-of-Custody
- 4. Data Summary Package (contains summary of all CLP or CLP-like Forms 1 through 15 per analytical fraction)
- 5. Analytical Fractions (e.g., VOA, SVOC, Metals, etc.)
  - a. Results and QC Summary (summary of all CLP or CLP like Forms 1 through 15 for a particular analytical fraction)
  - b. Raw Sample Data (includes all sample dilutions, sample re-analyses, QC samples, etc.)
  - c. Calibration Data (includes all initial and continuing calibrations and initial calibration verification)
  - d. Miscellaneous (includes extraction/preparation forms, percent solids determination, IDLs, MDLs, etc.)

Each of the above sections should be bookmarked in the PDF for easy access.

#### Summary Form Requirements for hardcopy and PDF data package deliverable for non-CLP Methods:

Second-source initial calibration verification summary forms are required, if applicable per the method or the DOD QSM. Also, the analytes associated with internal standards must be identified.

The following summary forms are required as part of the data package deliverable for SW-846 6020/6010/7000 series for metals:

**Results Report** - must present all information presented on CLP FORM I (ILM05.4).

**Initial and Continuing Calibration Summary** - must present all information presented on CLP FORM 2A (ILM05.4). **Low-Level Calibration Standard Summary** – if applicable, must present all information presented on CLP FORM 2B (ILM05.4).

Blanks - must present all information presented on CLP FORM 3 (ILM05.4).

ICP Interference Check Sample Summary - must present all information presented on CLP FORM 4 (ILM05.4).

Matrix Spike Summary - must present all information presented on CLP FORM 5A (ILM05.4).

Post Digestion Spike - must present all information presented on CLP FORM 5B (ILM05.4).

Lab Duplicate Results - must present all information presented on CLP FORM 6 (ILM05.4).

LCS Summary - must present all information presented on CLP FORM 7 (ILM05.4).

**MSA Summary (Method of Standard Addition)** – if applicable, must present all information presented on CLP FORM 8 (ILM04.1).

ICP Serial Dilution Summary - must present all information presented on CLP FORM 8 (ILM05.4).

**Detection Limits** - must present all information presented on CLP FORM 9 (ILM05.4).

Linear Range – must present all information presented on CLP FORM 11 (ILM05.4).

Internal Standard Association (ICP-MS) –must present all information presented on CLP FORM 11 (ISM01.2). Alternatively, the laboratory may provide the information in the Narrative.

Prep Log - must present all information presented on CLP FORM 12 (ILM05.4).

Analysis Run Log - must present all information presented on CLP FORM 13 (ILM05.4).

**ICP-MS Tune** – must present all information presented on CLP FORM 14 (ISM01.2). Laboratory must also document the number of tune analysis integrations on this form or in the Narrative.

**ICP/MS Internal Standard Relative Intensity Summary** - must present all information presented on CLP FORM 15 (ILM05.4).

<u>Also must include</u>: Instrument Calibration Records, Chain-of-Custody Forms, and Case Narrative. The Narrative, forms, or raw data must indicate the number of replicate integrations for ICP-MS sample analysis.

Summary Forms for SW-846, 8260 and 8270 (i.e., any SW-846 GC/MS analysis of Volatile and Semivolatile Organic Compounds) should be presented in a CLP-Like format. The following Summary Forms must be included:

Result Summary One Sample per summary page.

Presentation of analytical results for both method blanks and environmental samples, date of collection, preparation, and analysis. Environmental samples should be identified with the field identification

numbers on the COCs.

Surrogate Recovery Form Present all information contained on CLP Form II.

Summary of Matrix Spike/Matrix Spike Duplicate

Recovery

Present all information contained on CLP Form III.

Instrument Performance Check Summary Form -

Mass Spec Tuning Form

Present all information Contained on CLP Form V.

Initial Calibration Summary Present all information contained CLP Form VI.

Continuing Calibration Summary Present All Information contained on CLP Form VII.

Internal Standard Area and Retention Time

Summary

Present all information contained CLP Form VIII.

### Summary Forms for GC analysis should be presented in a CLP-Like format. The following Summary Forms must be included:

Result Summary One Sample per summary page.

Presentation of analytical results for both method blanks and environmental samples, date of collection, preparation, and analysis. Environmental samples should be identified

with the field identification numbers on the COCs.

Surrogate Recovery Form Present all information contained on CLP Form II for both

Analytical Columns.

Summary of Matrix Spike/Matrix Spike

**Duplicate Recovery** 

Present all information contained on CLP Form III.

Summary of Initial Calibration of Single

Component Analytes

Present all information contained on CLP Form VI-PEST-2.

Summary of Calibration Verification Present all information contained on CLP Form VII-PEST-1

and Form VII-PEST-2.

Analytical Sequence Present all information contained on CLP Form VIII-PEST.

**Identification Summary** 

For Single Component Analytes and for Multiple Component Analytes

Present all information contained on CLP Form X PEST 1

and 2.

ATTACHMENT C
ELECTRONIC DATA DELIVERABLE REQUIREMENTS

#### **ELECTRONIC DATA FORMAT REQUIREMENTS**

#### 1.0 INTRODUCTION

The laboratory is to submit text-based tab delimited EDD files for each SDG using Tetra Tech's laboratory data checker explained below. The files must be in the format specified in this Attachment. Additional information such as laboratory name, project name, fractions included, project number, site name/number, laboratory contact person and any specific comments related to the EDD should be included in the comments section of the EDD Submittal page.

The RESULT for nondetects should be populated with the project-specific sample quantitation reporting limits (i.e., either the sample quantitation limit or method detection limit, as specified in Section 3.0 of this scope of work. Any corrections made to the hardcopy data must also be made to the electronic file. Appropriate qualifiers as identified by the analytical protocol must also be designated; laboratory QC non-compliance codes are not to be depicted.

Tetra Tech's electronic EDD format follows the ADAPT structure and requires the A1 and A3 files. The A2 file is only required if the project is using ADAPT; and, for non-ADAPT EDD submittals the A2 file may be omitted. The EDD consists of separate, tab-delimited ASCII text files. Each file corresponds to a database table. The tables are identified as the Analytical Results Table (A1) and Sample Analysis Table (A3). A separate set of text files must be created and submitted for each sample delivery group (SDG). The files must be identified to correspond to the (A1) table and the (A3) table. The file naming convention is: the Sample Delivery Group (SDG) followed by the table identifier (A1 or A3), followed by the ".txt" extension. The file names must not contain spaces or special characters. For example, the EDD file names for a laboratory-reporting batch identified as SDG001 would be as follows:

SDG001A1.txt SDG001A3.txt

On certain projects Tetra Tech will utilize the ADAPT Electronic Data Validation software, which will require the laboratory to use the ADAPT electronic data deliverable checker software prior to submitting the files through Tetra Tech's laboratory data checker (this will be clearly specified in the Tetra Tech laboratory statement of work). The ADAPT checker software can be downloaded from Laboratory Data Consultants' web site: http://www.lab-data.com. For projects which Tetra Tech is using the ADAPT software, Tetra Tech will provide the laboratory with the project library. The laboratory is not permitted to modify the project library. ADAPT projects will require the laboratory to export all three checked files (A1, A2, and A3) from the ADAPT software and submit them through Tetra Tech's laboratory data checker. ADAPT error logs generated must be included with the electronic PDF data validation package and cannot be submitted through the laboratory data checker.

The values reported in the EDD text files must agree exactly with the final values reported on the PDF data package sample result summaries. The details of file naming conventions, data structure and data checker use are discussed below.

### **Analytical Results Table (A1 File)**

The Analytical Results table contains analytical results and related information for target analytes in field samples and associated laboratory quality control samples (excluding calibrations and tunes). Field samples and laboratory method blanks must report a result record for each analyte reported within a method. Laboratory control samples (LCS and LCSD) and matrix spike samples (MS and MSD) must report a result record for every analyte specified as a spiked analyte in the laboratory statement of work. Table A1 in this document lists the field names and data type descriptions for the Analytical Results Table (A1).

### Lab Instrument Table (A2 File)

A2 file is only required if the project is using ADAPT. In all other EDD submittals, the A2 file may be omitted. Laboratories should refer to the ADAPT User Guide for populating the A2 Table.

### Sample Analysis Table (A3 File)

The Sample Analysis table contains information specific to field environmental samples and laboratory quality control analyses (excluding calibrations and tunes). A sample record must exist for each sample/method/matrix/analysis type combination. Table A3 in this document lists the field names and data type descriptions for the Sample Analysis Table (A3).

All electronic data deliverables are due within the same time established for the associated hardcopy data packages.

In addition, the laboratory QC officer must read and sign a copy of the Quality Assurance Review Form displayed on the next page of this Attachment. Electronic deliverables are not considered to be complete without the accompanying Quality Assurance Review Form.

I	ables have be a. The enclores), completer to correct enclores	sed electronic f ness and format.	viewed and are iles have been The laboratory	in agreement reviewed for will be respon	t with the accuracy nsible for
( )		any questions or	problems with	the enclosed	electronic
deliverables.					
Signature:		Titlo.		Date:	

### 2.0 EDD Field Properties

Tables A1 and A3 in this document specify the EDD field properties. Laboratories should refer to the ADAPT User Guide for populating the A2 Table. These include the field name, sequence order, field description, data type/length and reporting requirement for each field. Fields in the EDD **must** be sequenced according to the order that they appear below in Tables A1 and A3. For example, in the Analytical Results table (A1), the field "ClientSampleID" will always be the first piece of information to start every new line of data (or database record), followed by the field "LabAnalysisRefMethodID", "AnalysisType", etc.

When creating an EDD as a text file, use the ASCII character set in a file of lines terminated by a carriage return and line feed. No extra characters are allowed at the end of a line, after the carriage return and line feed. Enclose each data value with double quotes (text qualifier) and separate each field value with a **tab** character (tab delimiter). Data fields with no information (null) may be represented by two consecutive tabs. For example, in the Sample Analysis table, since the "Collected", "ShippingBatchID", and "Temperature" fields do not apply to laboratory generated QA/QC samples, the record for a Laboratory Control Sample by Method 8270C would be entered as follows. Note that the first two fields ("ProjectNumber" and "ProjectName") are omitted in this example.

..."LCSW100598" "AQ" "LCSW100598" "LCS" "8270C",...etc. If a field is populated with less than the maximum allowed number of characters, do not pad the values with leading or trailing spaces. In the above example, although the "MatrixID" field can accommodate up to 10 characters, only 2 characters were entered in this field. **Do not include the delimiter (tab character) within any of the field values.** Example EDD files may be downloaded from the LEDD Checker application.

An example database shall be sent for review prior to the first electronic deliverable in the required .txt format. The example file will be examined for completeness and comments will be sent to the laboratory. Any questions regarding the electronic deliverable should be directed to <a href="mailto:LabSupport@tetratech.com">LabSupport@tetratech.com</a>

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

Contains laboratory test results and related information for field and QC samples (excluding instrument calibrations) on an analyte level for environmental chemistry including radiochemistry

radioonemistry				
Field Name	Field Name Description	Field Type	Field Length	Required Value
ClientSampleID	Client or contractor's identifier for a field sample	Text	25	X
	If a sample is analyzed as a laboratory duplicate, matrix spike, or matrix spike duplicate, append suffixes DUP, MS and MSD respectively to the Client Sample ID			

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiochemistry			1	
=		Field	Field	Required
Field Name	with no intervening spaces or hyphens (i.e. MW01DUP, MW01MS, and MW01MSD). For Method Blanks, LCS, and LCSD enter the unique LaboratorySampleID into this field.  Do not append suffixes to the ClientSampleID for dilutions, reanalyses, or re-extracts (the AnalysisType field is used for this distinction). For example, MW01DL and MW01RE are not allowed.  Parent sample records must exist for each MS and MSD. If an MS/MSD is shared between two EDDs, records for the MS/MSD and its parent sample must exist in the Analytical Results table for both EDDs.	Type	Length	Value
LabAnalysisRefMetho dID	Laboratory reference method ID. The method ID may be an EPA Method number or a Lab Identifier for a method such as a SOP Number, however; method ID is specified by the project. The method ID must be entered into the standard list.	Text	25	X
AnalysisType	Defines the analysis type (i.e., Dilution, Reanalysis, etc.). This field provides distinction for sample result records when multiple analyses are submitted for the same sample, method, and matrix; for example dilutions, re-analyses, and re-extracts.	Text	10	X
LabSampleID	Laboratory tracking number for field samples and lab generated QC samples such as method blank, LCS, and LCSD. There are no restrictions for the LabSampleID except for field length and that the LabSampleID must be distinct for a given field sample or lab QC sample and method.	Text	25	X

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiochemistry		Field	Field	Deguired
Field Name	Field Name Description	Field Type	Field Length	Required Value
	Suffixes may be applied to the LabSampleID to designate dilutions, reanalysis, etc.			
LabID	Identification of the laboratory performing the analyses.	Text	7	X
ClientAnalyteID	CAS Number or unique client identifier for an analyte or isotope.  If a CAS Number is not available, use a unique identifier provided by the client or contractor. The ClientAnalyteID for a particular target analyte or isotope should be specified by the project and must exist in the standard value tables for Analytes.  For the LCS, LCSD, MS, and MSD, it is only necessary to report the compounds designated as spikes in the library (and surrogates for organic methods.)  For TICs from GC/MS analyses, enter the retention time in decimal minutes as the Client Analyte ID.	Text	12	X
AnalyteName	Chemical name for the analyte or isotope. The project specifies how an analyte or isotope is named. The analyte name must be associated to a ClientAnalyteID in the standard values table for Analytes (excluding compounds designated as TIC's).	Text	60	X
Result	Result value for the analyte or isotope.  Entries must be numeric. For non-	Numeric (1)	20(6)	X

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiocnemistry				<b>.</b>
Field Name	Field Name Description	Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
	detects of target analytes or isotopes and spikes, do not enter "ND" or "0". Do not leave this field blank. If an analyte or spike was not detected, enter the associated value specified in Section 3.0 of this scope of work (e.g., LOD, SQL, PQL, etc.), corrected for dilution and percent moisture as applicable. Do not enter "0". A "0" result may be acceptable for surrogate or internal standard percent recoveries; however, it should not be reported for any target compound.			
ResultUnits	The units defining how the values in the Result, DetectionLimit, and ReportingLimit fields are expressed. For radiochemistry this also includes how the value in the Error field is expressed.	Text	10	X
LabQualifiers	A string of single letter result qualifiers assigned by the lab based on client-defined rules and values.  The "U" Lab Qualifier must be entered for all non-detects. Other pertinent lab qualifiers may be entered with the "U" qualifier. Order is insignificant. Lab qualifiers other than those listed in the standard values table may be used. If so, these must be added to the standard value table in the application.	Text	7	Q
DetectionLimit	For radiochemistry methods, the minimum detectable activity for the isotope being measured.  For all other methods: The minimum	Numeric <sup>(1)</sup>	10(6)	X

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiochemistry			1	,
_		Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
	detection limit value for the analyte being measured.  For surrogates, internal standards, etc. where detection limits are not applicable use the value -99.			
DetectionLimitType	Specifies the type of detection limit (i.e., MDA, MDL, IDL, etc.).  If -99 is specified in the DetectionLimit field us the value NA.	Text	10	X
RetentionTime or Error	For radiochemistry methods only, enter the 2 Sigma Counting Errors. The units for error are entered in the ResultUnits field.  For GC/MS methods only, enter the time expressed in decimal minutes between injection and detection for GC/MS TICs only  For target analytes in all other methods, leave this field blank. Note: GC retention times are not evaluated at this time.	Text	5	T
AnalyteType	Defines the type of result, such as tracer, surrogate, spike, or target compound.	Text	7	X
PercentRecovery	For radiochemistry methods: The tracer yield, if applicable.  For all other analytical methods: The percent recovery value of a spiked compound or surrogate.  If the spike or surrogate was not recovered because of dilution, enter "DIL". If a spike or surrogate was not recovered because of matrix interference, enter "INT". If a spike or	Numeric <sup>(1)</sup>	5(3)	X

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiocnemistry		F:ald	Field.	Dearrined
Field Name	Field Name Description	Field Type	Field Length	Required Value
	surrogate was not recovered because it was not added to the sample, enter "NS".			
RelativePercentDiffere nce	The relative percent difference (RPD) of two QC results, such as MS/MSD, LCS/LCSD, and Laboratory Duplicates. Report RPD in Laboratory Duplicate, LCSD, and MSD records only.	Numeric (1)	5(3)	X
ReportingLimit	Reporting limit value for the measured analyte or isotope Factor in the dilution factor and percent moisture correction, if applicable. The Reporting Limit for each analyte and matrix in a given method is specified in the project library or QAPP.  For surrogates, internal standards, etc. where reporting limits are not applicable use the value -99.	Numeric (1)	10(6)	X
ReportingLimitType	Specifies the type of reporting limit (i.e., CRQL, PQL, SQL, RDL, etc). The Reporting Limit Type for each method and matrix is specified in the project library or QAPP.  If -99 is specified in the ReportingLimit field us the value NA.	Text	10	X
ReportableResult	This field indicates whether or not the laboratory chooses an individual analyte or isotope result as reportable. Enter "YES" if the result is reportable. Enter "NO" if the result is not reportable.  If only one analysis is submitted for a particular sample and method, enter "YES"	Text	3	X

Table A1 Field Descriptions for the Analytical Results Table (Table A1)

radiochemistry				
		Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
	for all target compounds (where Analyte Type = TRG). For GC/MS methods enter yes for tentatively identified compounds (where Analyte Type = TIC).			
	If two or more analyses are submitted for a particular sample and method (i.e. initial analysis, reanalysis and/or dilutions), enter "YES" from only one of the analyses for each target compound. For example: a sample was run a second time at dilution because benzene exceeded the calibration range in the initial, undiluted analysis. All target analytes are reported in each analysis. For the initial analysis, (Analysis Type = RES), enter "NO" for benzene and enter "YES" for all other compounds. For the diluted analysis (Analysis Type = DL), enter "YES" for benzene and enter "NO" for all other compounds.			
	For TICs (Analyte Type = TIC), if more than one analysis is submitted for a particular sample and method, choose only one of the analyses where Reportable Result = YES for all TICs. For example, a sample was run a second time because one or more target compounds exceeded the calibration range in the undiluted analysis. Choose a particular analysis and enter "YES" for all TICS. In the other analysis enter "NO" for all TICs.			
	Note that it is not necessary to report the full target analyte list for the initial result, dilution, re-analysis, or re-extraction. However, each target analyte must be reported YES once and once only in the case of multiple analyses for a given sample, method, and			

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

radiocnemistry		Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
	matrix. In the case of organics, all surrogates must be reported for all analyses submitted for a given sample, method, and, matrix.			
SpkConcnAdded	The spike added. This value must be reported in the same units as the result. Where (SA) in the following equation: % Recovery = (SSA-SC)/SA x 100% where: SSA is the spiked sample concentration (amount) after spiking. SC is the sample concentration (amount) before spiking. SA is the the expected increase in sample concentration (amount) as a result of spiking. This value must incorporate all correction factors such as dilution factor and moisture content that are applied to the spiked sample when computing the spiked sample concentration or amount. Enter -99 where no spike was added.	Numeric (1)	10(6)	X
SpkParentSampleID	The sampleID of a sample (often called the original sample) that receives a spike aliquot to form a spiked sample such as a matrix spike. This is not the same as the ID of the spiked sample (such as a matrix spike) after spiking.  The result for SpkParentSampleID and the result (i.e., SpkConcnAdded) for the spiked sample are used to compute percent recovery of the analyte.	Text	25	X
SamplePrepInitial	The initial sample preparation volume in liters (L) for aqueous samples or grams (g) for solid samples.	Numeric <sup>(1)</sup>	20(6)	

Table A1
Field Descriptions for the Analytical Results Table (Table A1)

Field Name	Field Name Description	Field Type	Field Length	Required Value
SamplePrepFinal	The final sample preparation volume in liters (L) for aqueous samples or grams (g) for solid samples.	Numeric (1)	20(6)	
LimitOfDetection	The smallest amount or concentration of a substance that must be present in a sample in order to be detected at a 99% confidence level. In other words, if a sample has a true concentration at the LOD, there is a minimum probability of 99% of reporting a "detection" (a measured value ≥ DL) and a 1% chance of reporting a non-detect (a false negative).	Numeric (1)	10(6)	N
Comment	Add any comments or additional information specific to the analyte test result data record.	Text	200	

- X Required field.
- Q Only required if laboratory has qualified the result.
- T Only required for tentatively identified compounds by GC/MS.
  - (1) Field Length indicates decimal precision in parenthesis. For example, 5(2) = a total width of 5 numbers including a maximum of 2 decimal places.

Field Description for the Sample Analysis (Table A3)

This table contains information related to analyses of field samples and laboratory QC samples (excluding calibrations and tunes) on a sample level for environmental chemical analyses including radiochemistry

	ennou y	Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
ProjectNumber	Project number assigned by the client.	Text	30	X
ProjectName	Project name assigned by the client.	Text	90	X
ClientSampleID	Client or contractor's identifier for a field sample	Text	25	X
	If a sample is analyzed as a laboratory duplicate, matrix spike, or matrix spike duplicate, append suffixes DUP, MS and MSD respectively to the Client Sample ID with no intervening spaces or hyphens (i.e. MW01DUP, MW01MS, and MW01MSD). For Laboratory QC samples (i.e. Method Blanks, LCS, and LCSD, etc.) enter the unique LaboratorySampleID into this field			
	Do not append suffixes to the ClientSampleID for dilutions, reanalyses, or re-extracts (the Analysis_Type field is used for this distinction). For example, MW01 <u>DL</u> and MW01 <u>RE</u> are not allowed			
	Parent sample records must exist for each MS and MSD. If an MS/MSD is shared between two EDDs, records for the MS/MSD and its parent sample must exist in the Sample Analysis table for both EDDs.			
Collected	Date and Time of sample collection. Refer to the date/time format at the end of this table.  Leave this field blank for Method Blank, LCS, and LCSD. For Collected values that are not applicable use the value of 00/00/0000 00:00.	Date/ Time	16*	X
MatrixID	Sample matrix (i.e., AQ, SO, etc.)	Text	10	X
LabSampleID	Laboratory tracking number for field samples	Text	25	Х

Table A3 Field Description for the Sample Analysis (Table A3)

This table contains information related to analyses of field samples and laboratory QC samples (excluding calibrations and tunes) on a sample level for environmental chemical analyses including radiochemistry

including radiochen	noti y	Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
	and lab generated QC samples such as method blank, LCS, and LCSD.  There are no restrictions for the LabSampleID except field length and that the LabSampleID must be unique for a given field sample or lab QC sample and method.			
QCType	This record identifies the type of quality control sample QC (i.e., Duplicate, LCS, Method Blank, MS, or MSD). For regular environmental samples, populate this field with "NM".	Text	10	X
ShippingBatchID	Unique identifier assigned to a cooler or shipping container used to transport client or field samples. Links all samples to a cooler or shipping container. No value is required for method blanks, LCS, and LCSD.	Text	25	Х
Temperature	Temperature (in centigrade degrees) of the sample as received.  The storage refrigerator or room temperature should be reported (in centigrade degrees) for laboratory QC samples (i.e. method blanks, laboratory control standards).  Use -99 if temperature is not available.  This field is not required for radiochemistry methods.	Numeric (1)	10(6)	X
LabAnalysisRefMeth odID	Laboratory reference method ID. The method ID may be an EPA Method number or laboratory identifier for a method such as a SOP number, however; values used for Laboratory Method IDs are specified by the	Text	25	X

Table A3

Field Description for the Sample Analysis (Table A3)

This table contains information related to analyses of field samples and laboratory QC samples (excluding calibrations and tunes) on a sample level for environmental chemical analyses including radiochemistry

Field Name	Field Name Description	Field Type	Field Length	Required Value
	project and must in the in standard value list for method IDs.			
PreparationType	Preparation Method Number (i.e., 3010A, 3510C, 3550C, 5030B, etc.)  For analytical procedures that do not have a specific preparation method number, use "Gen Prep".	Text	25	X
AnalysisType	Defines the type of analysis such as initial analysis, dilution, re-analysis, etc. This field provides distinction for sample records when multiple analyses are submitted for the same sample, method, and matrix, for example: dilutions, re-analyses, and re-extracts.	Text	10	X
Prepared	Refer to the date/time format at the end of this table. If no sample preparation is involved enter the analysis date and time in this field. Refer to the date/time format at the end of this table.	Date/ Time	16*	X
Analyzed	Date and time of sample analysis. Refer to the date and time format at the end of this table. For Analyzed values that are not applicable use the value of 00/00/0000 00:00.	Date/ Time	16*	X
LabID	Identification of the laboratory performing the analysis.	Text	7	X
QCLevel	The level of laboratory QC associated with the analysis reported in the EDD. If only the Analytical Results Table (A1) and the Sample Analysis Table (A3) information are submitted for the sample, enter "COA". If the Laboratory Instrument Table (A2) information is also submitted for the sample, enter "COCAL"	Text	6	X

Table A3 Field Description for the Sample Analysis (Table A3)

This table contains information related to analyses of field samples and laboratory QC samples (excluding calibrations and tunes) on a sample level for environmental chemical analyses including radiochemistry

ncluding radiochemistry Field Field Require				
Field Name	Field Name Description	Type	Length	Required Value
ResultBasis	Indicates whether results associated with this sample records are reported as wet or percent moisture corrected. Enter "WET" if results are not corrected for percent moisture. Enter "DRY" if percent moisture correction is applied to results. For aqueous samples, enter "WET". For other matrices where basis is not applicable enter "NA"		3	X
TotalOrDissolved	This field indicates if the results related to this sample record are reported as a total or dissolved fraction. If not applicable please report "NA"	Text	3	Х
Dilution	Dilution of the sample aliquot. Enter "1" for method blanks, LCS, and LCSD, or if the field samples was analyzed without dilution.	Numeric (1)	10(6)	X
HandlingType	Indicates the type of leaching procedure, if applicable (i.e., SPLP, TCLP, WET).  Enter "NA" if the sample analysis was not performed on a leachate.	Text	10	X
HandlingBatch	Unique laboratory identifier for a batch of samples prepared together in a leaching procedure (i.e., SPLP, TCLP, or WET preparation). The HandlingBatch links samples with leaching blanks.  Enter "NA" if the sample analysis was not performed on a leachate.	Text	12	X
LeachateDate	Date and time of leaching procedure (i.e., date for SPLP, TCLP, or WET preparation). Refer to the date and time format at the end of this table.	Date /Time	16*	X

Table A3

Field Description for the Sample Analysis (Table A3)

This table contains information related to analyses of field samples and laboratory QC samples (excluding calibrations and tunes) on a sample level for environmental chemical analyses including radiochemistry

Field Name		Field	Field	Required
rieid Name	Field Name Description  . For Analyzed values that are not applicable use the value of 00/00/0000 00:00	Туре	Length	Value
Percent_Moisture	For soil and sediment samples, enter the percent of sample composed of water. For aqueous samples enter "100". For other matrices where Percent_Moisture is not applicable use a value of -99	Numeric (1)	10(6)	X
MethodBatch	Unique laboratory identifier for a batch of samples of similar matrices analyzed by one method and treated as a group for matrix spike, matrix spike duplicate, or laboratory duplicate association  The method batch links the matrix spike and/or matrix spike duplicate or laboratory duplicates to associated samples. Note the MethodBatch association may coincide with the PreparationBatch association. The MethodBatch is specifically used to link the MS/MSD and/or DUP to associated samples.		12	X
PreparationBatch	Unique laboratory identifier for a batch of samples prepared together for analysis by one method and treated as a group for method blank, LCS and LCSD association.  The PreparationBatch links method blanks and laboratory control samples (blank spikes) to associated samples. Note, the PreparationBatch association may coincide with the MethodBatch association but the PreparationBatch specifically links the Method Blank and LCS to associated samples.	Text	12	X
RunBatch	For all other methods the RunBatch is the unique identifier for a batch of analyses performed on one instrument under the control	Text	12	X

Table A3 Field Description for the Sample Analysis (Table A3)

This table contains information related to analyses of field samples and laboratory QC samples (excluding calibrations and tunes) on a sample level for environmental chemical analyses including radiochemistry

including radiochemistry					
		Field	Field	Required	
Field Name	Field Name Description	Type	Length	Value	
	of one initial calibration and initial calibration verification. The RunBatch links both the initial calibration and initial calibration verification to subsequently analyzed and associated continuing calibrations, field samples, and QC analyses. For GC/MS methods, the RunBatch also links a BFB or DFTPP tune. A distinct RunBatch must used with every new initial calibration within a method  The value entered in this field links a particular sample/method/analysis type record to a set of associated initial calibration and initial calibration verification records from Table A2.  If Table A2 is not submitted enter a value of 'NA" in this field.				
AnalysisBatch	For radiochemistry methods leave this field blank.  For all other methods the AnalysisBatch is the unique identifier for a batch of analyses performed on one instrument and under the control of a continuing calibration or continuing calibration verification. The AnalysisBatch links the continuing calibration or calibration verification to subsequently analyzed and associated field sample and QC analyses. For GC/MS methods, the AnalysisBatch also links the BFB or DFTPP tune. A distinct AnalysisBatch must be used with every new continuing calibration or continuing calibration verification within a method  The value entered in this field links a particular sample/method/analysis type record to a set of		12	X	

### Table A3 Field Description for the Sample Analysis (Table A3)

This table contains information related to analyses of field samples and laboratory QC samples (excluding calibrations and tunes) on a sample level for environmental chemical analyses

including radiochemistry

increaning radiocites		Field	Field	Required
Field Name	Field Name Description	Type	Length	Value
	associated continuing calibration records in the Laboratory Instrument table.			
LabReportingBatch	Unique laboratory identifier for the EDD. This is equivalent to the sample delivery group, lab work number, login ID, etc. The LabReportingBatch links all records in the EDD reported as one group. The value entered in this field must be the same in all records.	Text	12	X
LabReceipt	Date and time the sample was received in the lab. A time value of 00:00 may be entered. Refer to the date/time format at the end of this table.	Date/ Time	16*	X
LabReported	Date and time hard copy reported delivered by the lab. A time value of 00:00 may be entered. Refer to the date/time format at the end of this table.		16*	X
Comment	Add any comments or additional information specific to the sample analysis data record.	Text	200	

C Only required for regular samples, duplicates and MS/MSDs.

X Required field.

### 3.0 Laboratory Data Checker

The Laboratory Data Checker is a web-based application that will review Laboratory Electronic Data Deliverables (LEDDs) for adherence to Tetra Tech's EDD format requirements. EDDs will be reviewed for elements such as missing data and/or columns of data, and compliance of the data within each column to the required data types/lengths.

<sup>(1)</sup> Field Length indicates decimal precision in parenthesis. For example, 5(2) = a total width of 5 numbers including a maximum of 2 decimal places.

<sup>\*</sup> Format Date and Time as MM/DD/YYYY hh:mm; where MM = two digit month, DD = two digit day, and YYYY = four digit year, hh = hour in 24 hour format, and mm = minutes.

Once an EDD passes through the checker with no errors, it must be submitted to Tetra Tech through the LEDD Checker application.

Access to the LEDD Checker application will be provided by an initial registration/approval process. An Information Systems Group (ISG) Administrator will approve requests for access. To access the site or begin the registration process, visit the ISG web site at <a href="http://isg.ttnus.com">http://isg.ttnus.com</a> and select the "Laboratory Checker" link on the left of the home page. Registered users may access the checker immediately by logging in to the system using theircredentials. New users must select the "Register" button and provide all of the requested information.

After completing all fields on the registration form, select the "Submit" button to complete the request process. Upon verification by an ISG Administrator, an email notification will be sent verifying the user ID, password and account status. Forgotten passwords may be retrieved by using the "Forgot password?" link on the login page. Note that the email address that was provided for registration or password retrieval is the user ID and must be a valid e-mail address.

The general process for submitting EDD files through the LEDD Checker involves a 3-stage process that includes an upload stage, an error checking stage and a submittal stage.

Log into the LEDD Checker by typing your login credentials and select the "Login" button. The LEDD Checker home page provides a general overview of the checker functionality and EDD file format requirements. At the bottom of the home page, example EDDs are provided that may be viewed or downloaded. To download the files, right click on the link and select "Save target as" from the menu. Each LEDD Checker page includes a navigation bar with links to return to the home page or continue the checking and submittal process. Users should **NOT** use the back or forward buttons on the browser, instead use the links provided in the application to navigate through the site.

Detailed information regarding EDD preparation, formatting requirements and text file naming conventions are provided in the Electronic Data Format Requirements Section of the Laboratory SOW.

Begin the upload stage by selecting the "Upload/Check Files" link on the home page. Follow the steps on the upload page starting with the selection of the laboratory name that corresponds to your organization. If your organization is not listed, contact <a href="mailto:LabSupport@tetratech.com">LabSupport@tetratech.com</a>, and provide a full description of your organization name, contact information and include "Laboratory Contractor ID Request" in the subject line. An ISG Administrator will respond to the request via e-mail.

Load the appropriate A1, A2, or A3 target EDD files by clicking the "Browse" button next to each data table input box. A file browser dialog will appear allowing files to be selected from a local or network drive. After the EDD files are loaded, click the "Upload" button to complete the upload stage. Note that each table may be uploaded and checked separately; however, a minimum of the A1 and A3 files are required in order to submit the EDDs.

If the file upload was successful, the checking page will immediately load. Begin the checking stage by selecting the "Check Files" button. The LEDD Checker will begin validating the EDD files for compliance. Depending on file size and network activity the validation process may take several minutes. The progress should be displayed in the information bar at the bottom of the browser window. **Do not** select the "Check Files" button again or otherwise use the browser during this process. Other applications may be used; however, note that the LEDD Checker may not sit idle for more than 30 minutes. If the time is exceeded a new session must be started in a new browser window.

Any errors will be processed and returned on the error page. The following general errors may be returned.

- Column count / table structure errors due to column header names being included, improper delimiter, extra tabs, extra or missing columns of data, spaces or other characters at the end of a row.
- Row and column value specific errors may occur for one or more reasons including: data truncation, invalid date / time format, invalid decimal precision or field width exceedance, or if a value is not in a list of valid values or expected range.

If column count / table structure errors are encountered, the LEDD Checker will return an error and stop the checking process.

The EDDs will not be processed any further until the column errors are resolved. Text fields are validated for truncation. Date / Time fields are validated for truncation and format compliance. Numeric decimal fields are validated for truncation, character type compliance and decimal precision. All required fields are validated for null values or empty text strings (i.e. spaces). The LEDD Checker will return a list of all errors in and include a reference to the row number on which the error occurred. Note that consecutive EDD files may be loaded and checked, and submitted while logged in. However, no data may be submitted until all EDD files have passed through the LEDD Checker without errors. The list of errors may be printed by selecting the "Print this Page" button from the checker error page.

If the EDD files pass with no errors, the submittal page will immediately load. To complete the submittal stage, include the following information in the comment and additional information area of the form: laboratory name, laboratory contact person, project name, project number, site name/number, fractions included and any specific comments related to the EDD. Select the "Submit Files" button to continue the submittal process.

The submittal stage is not considered complete until a unique ticket key reference is returned in the browser window. The ticket key reference must be printed for record of submission and future reference. In addition, a copy of the ticket key reference must be included in the PDF data package.

## APPENDIX F LABORATORY CERTIFICATION AND SOPS





### Certificate of Accreditation

ISO/IEC 17025:2005

Certificate Number L2223

### Katahdin Analytical Services, Inc.

600 Technology Way Scarborough ME 04074

has met the requirements set forth in L-A-B's policies and procedures, all requirements of ISO/IEC 17025:2005 "General Requirements for the competence of Testing and Calibration Laboratories" and the U.S. Department of Defense Environmental Laboratory Accreditation Program (DoD ELAP).\*

The accredited lab has demonstrated technical competence to a defined "Scope of Accreditation" and the operation of a laboratory quality management system (refer to joint ISO-ILAC-IAF Communiqué dated 8 January 2009).

Accreditation valid through: February 1, 2016

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R. Douglas Leonard, Jr., President, COO Laboratory Accreditation Bureau Presented the 1st of February 2013



# Scope of Accreditation For Katahdin Analytical Services, Inc.

600 Technology Way Scarborough, ME 04074 Leslie Dimond 207-874-2400

In recognition of a successful assessment to ISO/IEC 17025:2005 and the requirements of the DoD Environmental Laboratory Accreditation Program (DoD ELAP) as detailed in the DoD Quality Systems Manual for Environmental Laboratories (DoD QSM v4.2) based on the National Environmental Laboratory Accreditation Conference Chapter 5 Quality Systems Standard (NELAC Voted Revision June 5, 2003), accreditation is granted to Katahdin Analytical Services to perform the following tests:

Accreditation granted through: February 1, 2016

### **Testing - Environmental**

Non-Potable Water			
Technology	Method	Analyte	
GC/ECD	EPA 8081B	2, 4`-DDD	
GC/ECD	EPA 8081B	2, 4`-DDE	
GC/ECD	EPA 8081B	2, 4`-DDT	
GC/ECD	EPA 608; EPA 8081B	4, 4`-DDD	
GC/ECD	EPA 608; EPA 8081B	4, 4`-DDE	
GC/ECD	EPA 608; EPA 8081B	4, 4`-DDT	
GC/ECD	EPA 608; EPA 8081B	Aldrin	
GC/ECD	EPA 608; EPA 8081B	alpha-BHC (alpha-Hexachlorocyclohexane)	
GC/ECD	EPA 8081B	Alpha-Chlordane	
GC/ECD	EPA 608; EPA 8081B	beta-BHC (beta-Hexachlorocyclohexane)	
GC/ECD	EPA 8081B	Cis-Nonaclor	
GC/ECD	EPA 608; EPA 8081B	Chlordane (tech.)	
GC/ECD	EPA 608; EPA 8081B	delta-BHC	
GC/ECD	EPA 608; EPA 8081B	Dieldrin	
GC/ECD	EPA 608; EPA 8081B	Endosulfan I	
GC/ECD	EPA 608; EPA 8081B	Endosulfan II	

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-Potable Water			
Technology	Method	Analyte	
GC/ECD	EPA 608; EPA 8081B	Endosulfan sulfate	
GC/ECD	EPA 608; EPA 8081B	Endrin	
GC/ECD	EPA 608; EPA 8081B	Endrin aldehyde	
GC/ECD	EPA 8081B	Endrin Ketone	
GC/ECD	EPA 8081B	gamma-BHC (Lindane gamma- Hex <mark>achloro</mark> cyclohexane)	
GC/ECD	EPA 8081B	gamma-Chlordane	
GC/ECD	EPA 608; EPA 8081B	Heptachlor	
GC/ECD	EPA 608; EPA 8081B	Heptachlor epoxide	
GC/ECD	EPA 8081B	Hexachlorobenzene	
GC/ECD	EPA 8081B	Methoxychlor	
GC/ECD	EPA 8081B	Mirex	
GC/ECD	EPA 8081B	Oxychlordane	
GC/ECD	EPA 608; EPA 8081B	Toxaphene (Chlorinated camphene)	
GC/ECD	EPA 8081B	trans-Nonachlor	
GC/ECD	EPA 608; EPA 8082A	Aroclor-1016 (PCB-1016)	
GC/ECD	EPA 608; EPA 8082A	Aroclor-1221 (PCB-1221)	
GC/ECD	EPA 608; EPA 8082A	Aroclor-1232 (PCB-1232)	
GC/ECD	EPA 608; EPA 8082A	Aroclor-1242 (PCB-1242)	
GC/ECD	EPA 608; EPA 8082A	Aroclor-1248 (PCB-1248)	
GC/ECD	EPA 608; EPA 8082A	Aroclor-1254 (PCB-1254)	
GC/ECD	EPA 608; EPA 8082A	Aroclor-1260 (PCB-1260)	
GC/ECD	EPA 8082A MOD	Aroclor-1262 (PCB-1262)	
GC/ECD	EPA 8082A MOD	Aroclor-1268 (PCB-1268)	
GC/ECD	EPA 8082A	2,2',3,3',4,4',5,5',6-Nonachlorobiphenyl (BZ 206)	
GC/ECD	EPA 8082A	2,2',3,3',4,4',5,6-Octachlorobiphenyl (BZ 195)	
GC/ECD	EPA 8082A	EPA 8082A 2,2',3,3',4,4',5-Heptachlorobiphenyl (BZ 170)	
GC/ECD	EPA 8082A	EPA 8082A 2,2',3,3',4,4'-Hexachlorobiphenyl (BZ 128)	
GC/ECD	EPA 8082A	2, 2', 3, 4, 4', 5, 5'-Heptachlorobiphenyl (BZ 180)	
GC/ECD	EPA 8082A	2, 2', 3, 4, 4', 5', 6-Heptachlorobiphenyl (BZ 183)	
GC/ECD	EPA 8082A	2, 2', 3, 4, 4', 5-Hexachlorobiphenyl (BZ 138)	
GC/ECD	EPA 8082A	2, 2', 3, 4, 4', 6, 6'-Heptachlorobiphenyl (BZ 184)	

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n-Potable Water		
Technology	Method	Analyte
GC/ECD	EPA 8082A	2, 2', 3, 4', 5, 5', 6-Heptachlorobiphenyl (BZ 187)
GC/ECD	EPA 8082A	2, 2', 3, 4, 5'-Pentachlorobiphenyl (BZ 87)
GC/ECD	EPA 8082A	2, 2', 3, 5'-Tetrachlorobiphenyl (BZ 44)
GC/ECD	EPA 8082A	2, 2', 4, 4', 5, 5'-Hexachlorobiphenyl (BZ 153)
GC/ECD	EPA 8082A	2, 2', 4, 5, 5'-Pentachlorobiphenyl (BZ 101)
GC/ECD	EPA 8082A	2, 2', 4, 5-Tetrachlorobiphenyl (BZ 48)
GC/ECD	EPA 8082A	2, 2', 4, 5'-Tetrachlorobiphenyl (BZ 49)
GC/ECD	EPA 8082A	2, 2', 5, 5'-Tetrachlorobiphenyl (BZ 52)
GC/ECD	EPA 8082A	2, 2', 5-Trichlorobiphenyl (BZ 18)
GC/ECD	EPA 8082A	2, 3, 3', 4, 4', 5-Hexachlorobiphenyl (BZ 156)
GC/ECD	EPA 8082A	2, 3, 3', 4, 4', 5'-Hexachlorobiphenyl (BZ 157)
GC/ECD	EPA 8082A	2, 3, 3', 4, 4'-Pentachlorobiphenyl (BZ 105)
GC/ECD	EPA 8082A	2, 3, 3', 4, 4', 5, 5'-Heptachlorobiphenyl (BZ 189
GC/ECD	EPA 8082A	2, 3', 4, 4', 5, 5'-Hexachlorobiphenyl (BZ 167)
GC/ECD	EPA 8082A	2, 3', 4, 4', 5-Pentachlorobiphenyl (BZ 118)
GC/ECD	EPA 8082A	2, 3', 4, 4',5'-Pentachlorobiphenyl (BZ 123)
GC/ECD	EPA 8082A	2, 3', 4, 4'-Tetrachlorobiphenyl (BZ 66)
GC/ECD	EPA 8082A	2, 3, 4, 4', 5-Pentachlorobiphenyl (BZ 114)
GC/ECD	EPA 8082A	2, 4, 4'-Trichlorobiphenyl (BZ 28)
GC/ECD	EPA 8082A	2, 4'-Dichlorobiphenyl (BZ 8)
GC/ECD	EPA 8082A	3, 3', 4, 4', 5, 5'-Hexachlorobiphenyl (BZ 169)
GC/ECD	EPA 8082A	3, 3', 4, 4', 5-Pentachlorobiphenyl (BZ 126)
GC/ECD	EPA 8082A	3, 3', 4, 4'-Tetrachlorobiphenyl (BZ 77)
GC/ECD	EPA 8082A	3, 4, 4', 5-Tetrachlorobiphenyl (BZ 81)
GC/ECD	EPA 8082A	Decachlorobiphenyl (BZ 209)
GC/ECD	EPA 8151A	2, 4, 5-T
GC/ECD	EPA 8151A	2, 4-D
GC/ECD	EPA 8151A	2, 4-DB
GC/ECD	EPA 8151A	Dalapon
GC/ECD	EPA 8151A	Dicamba
GC/ECD	EPA 8151A	Dichloroprop

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on-Potable Water		
Technology	Method	Analyte
GC/ECD	EPA 8151A	Dinoseb
GC/ECD	EPA 8151A	MCPA
GC/ECD	EPA 8151A	МСРР
GC/ECD	EPA 8151A	Pentachlorophenol
GC/ECD	EPA 8151A	Silvex (2, 4, 5-TP)
GC/FID	EPA 8015C/D MOI	Diesel range organics (DRO)
GC/FID	EPA 8015C/D MOI	Total Petroleum Hydrocarbon (TPH)
GC/FID	EPA 8015C/D MOI	Gasoline range organics (GRO)
GC/FID/PID	MA DEP VPH	Volatile Organic Hydrocarbons
GC/FID	MA DEP EPH	Extractable Petroleum Hydrocarbons
GC/FID	CT ETPH	Total Petroleum Hydrocarbons
GC/FID	TNRCC Method 100	Total Petroleum Hydrocarbons
GC/FID	FL-PRO	Petroleum Range Organics
GC/ECD	EPA 8011; EPA 504	1, 2-Dibromoethane (EDB)
GC/ECD	EPA 8011; EPA 504	1, 2-Dibromo-3-chloropropane
GC/FID	RSK-175	Methane Ethane Ethene
GC/MS	EPA 8260B/C; EPA 52	24.2 1, 1, 1, 2-Tetrachloroethane
GC/MS	EPA 624; EPA 8260B/C 524.2	EPA 1, 1, 1-Trichloroethane
GC/MS	EPA 624; 8260B/C EPA 524.2	1, 1, 2, 2-Tetrachloroethane
GC/MS	EPA 8260B/C	1,1,2-Trichloro-1,2,2-trifluoroethane
GC/MS	EPA 624; EPA 8260B/C 524.2	1, 1, 2-1 richioroethane
GC/MS	EPA 624; EPA 8260B/C 524.2	1, 1-Dichloroethane
GC/MS	EPA 624; EPA 8260B/C 524.2	EPA 1, 1-Dichloroethene
GC/MS	EPA 8260B/C; EPA 52	1, 1-Dichloropropene
GC/MS	EPA 8260B/C; EPA 52	1, 2, 3-Trichlorobenzene
GC/MS	EPA 8260B/C; EPA 52	1, 2, 3-Trichloropropane
GC/MS	EPA 8260B/C	1,2,3-Trimethylbenzene
GC/MS	EPA 8260B/C; EPA 52	24.2 1, 2, 4-Trichlorobenzene
GC/MS	EPA 8260B/C; EPA 52	24.2 1, 2, 4-Trimethylbenzene

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n-Potable Water		
Technology	Method	Analyte
GC/MS	EPA 8260B/C; EPA 524.2	1, 2-Dibromo-3-chloropropane
GC/MS	EPA 8260B/C; EPA 524.2	1, 2-Dibromoethane (EDB)
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	1, 2-Dichlorobenzene
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	1, 2-Dichloroethane
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	1, 2-Dichloropropane
GC/MS	EPA 8260B/C	1,3,5-Trichlorobenzene
GC/MS	EPA 8260B/C; EPA 524.2	1, 3, 5-Trimethylbenzene
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	1, 3-Dichlorobenzene
GC/MS	EPA 8260B/C; EPA 524.2	1, 3-Dichloropropane
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	1, 4-Dichlorobenzene
GC/MS	EPA 8260B/C	1, 4-Dioxane
GC/MS	EPA 8260B/C	1-Chlorohexane
GC/MS	EPA 8260B/C; EPA 524.2	2, 2-Dichloropropane
GC/MS	EPA 8260B/C; EPA 524.2	2-Butanone
GC/MS	EPA 624; EPA 8260B/C	2-Chloroethyl vinyl ether
GC/MS	EPA 8260B/C; EPA 524.2	2-Chlorotoluene
GC/MS	EPA 8260B/C; EPA 524.2	2-Hexanone
GC/MS	EPA 8260B/C; EPA 524.2	4-Chlorotoluene
GC/MS	EPA 8260B/C; EPA 524.2	4-Methyl-2-pentanone
GC/MS	EPA 8260B/C; EPA 524.2	Acetone
GC/MS	EPA 8260B/C	Acetonitrile
GC/MS	EPA 624; EPA 8260B/C	Acrolein
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Acrylonitrile
GC/MS	EPA 8260B/C; EPA 524.2	Allyl chloride
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Benzene
GC/MS	EPA 8260B/C	Benzyl chloride
GC/MS	EPA 8260B/C; EPA 524.2	Bromobenzene
GC/MS	EPA 8260B/C; EPA 524.2	Bromochloromethane

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on-Potable Water		
Technology	Method	Analyte
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Bromodichloromethane
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Bromoform
GC/MS	EPA 8260B/C; EPA 524.2	Carbon disulfide
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Carbon tetrachloride
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Chlorobenzene
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Chloroethane
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Chloroform
GC/MS	EPA 8260B/C	Chloroprene
GC/MS	EPA 8260B/C; EPA 524.2	cis-1, 2-Dichloroethene
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	cis-1, 3-Dichloropropene
GC/MS	EPA 8260B/C	Cis-1,4-Dichloro-2-butene
GC/MS	EPA 8260B/C	Cyclohexane
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Dibromochloromethane
GC/MS	EPA 8260B/C; EPA 524.2	Dibromomethane
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Dichlorodifluoromethane
GC/MS	EPA 8260B/C; EPA 524.2	Diethyl ether
GC/MS	EPA 8260B/C	Di-isopropylether
GC/MS	EPA 8260B/C; EPA 524.2	Ethyl methacrylate
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Ethylbenzene
GC/MS	EPA 8260B/C	Ethyl-t-butylether
GC/MS	EPA 8260B/C; EPA 524.2	Hexachlorobutadiene
GC/MS	EPA 8260B/C	Iodomethane
GC/MS	EPA 8260B/C	Isobutyl alcohol
GC/MS	EPA 8260B/C	Isopropyl alcohol
GC/MS	EPA 8260B/C; EPA 524.2	Isopropyl benzene
GC/MS	EPA 8260B/C; EPA 524.2	m p-xylenes
GC/MS	EPA 8260B/C	Methyl acetate

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on-Potable Water		
Technology	Method	Analyte
GC/MS	EPA 8260B/C; EPA 524.2	Methacrylonitrile
GC/MS	EPA 624 / 8260B,C	Methyl bromide (Bromomethane)
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Methyl chloride (Chloromethane)
GC/MS	EPA 8260B/C; EPA 524.2	Methyl methacrylate
GC/MS	EPA 8260B/C; EPA 524.2	Methyl tert-butyl ether
GC/MS	EPA 8260B/C	Methylcyclohexane
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Methylene chloride
GC/MS	EPA 8260B/C; EPA 524.2	Naphthalene
GC/MS	EPA 8260B/C; EPA 524.2	n-Butylbenzene
GC/MS	EPA 8260B/C; EPA 524.2	n-Propylbenzene
GC/MS	EPA 8260B/C; EPA 524.2	o-Xylene
GC/MS	EPA 8260B/C	Pentachloroethane
GC/MS	EPA 8260B/C; EPA 524.2	p-Isopropyltoluene
GC/MS	EPA 8260B/C; EPA 524.2	Propionitrile
GC/MS	EPA 8260B/C; EPA 524.2	sec-butylbenzene
GC/MS	EPA 8260B/C; EPA 524.2	Styrene
GC/MS	EPA 8260B/C	t-Amylmethylether
GC/MS	EPA 8260B/C; EPA 524.2	tert-Butyl alcohol
GC/MS	EPA 8260B/C	tert-Butylbenzene
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Tetrachloroethene (Perchloroethylene)
GC/MS	EPA 8260B/C; EPA 524.2	Tetrahydrofuran
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Toluene
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	trans-1, 2-Dichloroethylene
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	trans-1, 3-Dichloropropylene
GC/MS	EPA 8260B/C; EPA 524.2	trans-1, 4-Dichloro-2-butuene
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Trichloroethene (Trichloroethylene)
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Trichlorofluoromethane
GC/MS	EPA 8260B/C	Vinyl acetate

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n-Potable Water		
Technology	Method	Analyte
GC/MS	EPA 624; EPA 8260B/C; EPA 524.2	Vinyl chloride
GC/MS	EPA 624 / 8260B,C	Xylene
GC/MS	EPA 8260B/C SIM	1,1,1,2-Tetrachloroethane
GC/MS	8260B, C SIM	1,1,1-Trichloroethane
GC/MS	EPA 8260B/C SIM	1,1,2,2-Tetrachloroethane
GC/MS	EPA 8260B/C SIM	1, 1, 2-Trichloroethane
GC/MS	EPA 8260B/C SIM	1,2,3-Trichloropropane
GC/MS	EPA 8260B/C SIM	1,1-Dichloroethane
GC/MS	EPA 8260B/C SIM	1,1-Dichloroethene
GC/MS	EPA 8260B/C SIM	1,2,4-Trichlorobenzene
GC/MS	EPA 8260B/C SIM	1,2,4-Trimethylbenzene
GC/MS	EPA 8260B/C SIM	1,2-Dibromo-3-chloropropane
GC/MS	EPA 8260B/C SIM	1,2-Dibromoethane
GC/MS	EPA 8260B/C SIM	1,2-Dichlorobenzene
GC/MS	EPA 8260B/C SIM	1,2-Dichloroethane
GC/MS	EPA 8260B/C SIM	1,2-Dichloropropane
GC/MS	EPA 8260B/C SIM	1,3-Dichlorobenzene
GC/MS	EPA 8260B/C SIM	1,3-Dichloropropane
GC/MS	EPA 8260B/C SIM	1,4-Dichlorobenzene
GC/MS	EPA 8260B/C SIM	2-Hexanone
GC/MS	EPA 8260B/C SIM	4-Methyl-2-pentanone
GC/MS	EPA 8260B/C SIM	Benzene
GC/MS	EPA 8260B/C SIM	Bromodichloromethane
GC/MS	EPA 8260B/C SIM	Carbon Tetrachloride
GC/MS	EPA 8260B/C SIM	Chloroform
GC/MS	EPA 8260B/C SIM	Chloromethane
GC/MS	EPA 8260B/C SIM	cis-1,2-Dichloroethene
GC/MS	EPA 8260B/C SIM	cis-1,3-Dichloropropene
GC/MS	EPA 8260B/C SIM	Dibromochloromethane
GC/MS	EPA 8260B/C SIM	Ethylbenzene
GC/MS	EPA 8260B/C SIM	Isopropylbenzene

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Гесhnology	Method	Analyte
GC/MS	EPA 8260B/C SIM	Hexachlorobutadiene
GC/MS	EPA 8260B/C SIM	Methylcyclohexane
GC/MS	EPA 8260B/C SIM	m,p-Xylene
GC/MS	EPA 8260B/C SIM	o-Xylene
GC/MS	EPA 8260B/C SIM	Tetrachloroethene
GC/MS	EPA 8260B/C SIM	trans-1,2-Dichloroethene
GC/MS	EPA 8260B/C SIM	Trans-1,3-Dichloropropene
GC/MS	EPA 8260B/C SIM	Trichloroethene
GC/MS	EPA 8260B/C SIM	Trichlorofluoromethane
GC/MS	EPA 8260B/C SIM	Vinyl Chloride
GC/MS	EPA 8260B/C SIM	Xylenes (total)
GC/MS	EPA 8270C/D	1, 2, 4, 5-Tetrachlorobenzene
GC/MS	EPA 625; EPA 8270C/D	1, 2, 4-Trichlorobenzene
GC/MS	EPA 625; EPA 8270C/D	1, 2-Dichlorobenzene
GC/MS	EPA 8270C/D	1, 2-Diphenylhydrazine
GC/MS	EPA 8270C/D	1, 3, 5-Trinitrobenzene
GC/MS	EPA 625; EPA 8270C/D	1, 3-Dichlorobenzene
GC/MS	EPA 8270C/D	1, 3-Dinitrobenzene
GC/MS	EPA 625; EPA 8270C/D	1, 4-Dichlorobenzene
GC/MS	EPA 8270C/D	1, 4-Dioxane
GC/MS	EPA 8270C/D	1, 4-Naphthoquinone
GC/MS	EPA 8270C/D	1, 4-Phenylenediamine
GC/MS	EPA 8270C/D	1-Chloronaphthalene
GC/MS	EPA 8270C/D	1-Methylnaphthalene
GC/MS	EPA 8270C/D	1-Naphthylamine
GC/MS	EPA 8270C/D	2, 3, 4, 6-Tetrachlorophenol
GC/MS	EPA 8270C/D	2, 4, 5-Trochlorophenol
GC/MS	EPA 625; EPA 8270C/D	2, 4, 6-Trichlorophenol
GC/MS	EPA 625; EPA 8270C/D	2, 4-Dichlorophenol
GC/MS	EPA 625; EPA 8270C/D	2, 4-Dimethylphenol
GC/MS	EPA 625; EPA 8270C/D	2, 4-Dinitrophenol

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-Potable Water			
Technology	Method	Analyte	
GC/MS	EPA 625; EPA 8270C/D	2, 4-Dinitrotoluene (2, 4-DNT)	
GC/MS	EPA 8270C/D	2, 6-Dichlorophenol	
GC/MS	EPA 625; EPA 8270C/D	2, 6-Dinitrotoluene (2, 6-DNT)	
GC/MS	EPA 8270C/D	2-Acetylaminofluorene	
GC/MS	EPA 625; EPA 8270C/D	2-Chloronaphthalene	
GC/MS	EPA 625; EPA 8270C/D	2-Chlorophenol	
GC/MS	EPA 625; EPA 8270C/D	2-Methyl-4 6-dinitrophenol	
GC/MS	EPA 8270C/D	2-Methylnaphthalene	
GC/MS	EPA 8270C/D	2-Methylphenol	
GC/MS	EPA 8270C/D	2-Naphthylamine	
GC/MS	EPA 8270C/D	2-Nitroaniline	
GC/MS	EPA 625; EPA 8270C/D	2-Nitrophenol	
GC/MS	EPA 8270C/D	2-Picoline	
GC/MS	EPA 8270C/D	3-Methylcholanthrene	
GC/MS	EPA 8270C/D	3-Nitroaniline	
GC/MS	EPA 8270C/D	4-Aminobiphenyl	
GC/MS	EPA 625; EPA 8270C/D	4-Bromophenyl phenyl ether	
GC/MS	EPA 625; EPA 8270C/D	4-Chloro-3-methylphenol	
GC/MS	EPA 8270C/D	4-Chloroaniline	
GC/MS	EPA 625; EPA 8270C/D	4-Chlorophenyl phenylether	
GC/MS	EPA 8270C/D	4-Dimethyl aminoazobenzene	
GC/MS	EPA 8270C/D	3, 4-Methylphenol	
GC/MS	EPA 8270C/D	4-Nitroaniline	
GC/MS	EPA 625; EPA 8270C/D	4-Nitrophenol	
GC/MS	EPA 8270C/D	4-Nitroquinoline-1-oxide	
GC/MS	EPA 8270C/D	5-Nitro-o-toluidine	
GC/MS	EPA 8270C/D	7, 12-Dimethylbenz(a)anthracene	
GC/MS	EPA 8270C/D	a a-Dimethylphenethylamine	
GC/MS	EPA 625; EPA 8270C/D	Acenaphthene	
GC/MS	EPA 625; EPA 8270C/D	Acenaphthylene	
GC/MS	EPA 8270C/D	Acetophenone	

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a-Potable Water		
Technology	Method	Analyte
GC/MS	EPA 8270C/D	Aniline
GC/MS	EPA 625; EPA 8270C/D	Anthracene
GC/MS	EPA 8270C/D	Aramite
GC/MS	EPA 8270C/D	Atrazine
GC/MS	EPA 8270C/D	Azobenzene
GC/MS	EPA 8270C/D	Benzaldehyde
GC/MS	EPA 625; EPA 8270C/D	Benzidine
GC/MS	EPA 625; EPA 8270C/D	Benzo(a)anthracene
GC/MS	EPA 625; EPA 8270C/D	Benzo(a)pyrene
GC/MS	EPA 625; EPA 8270C/D	Benzo(b)fluoranthene
GC/MS	EPA 625; EPA 8270C/D	Benzo(g h i)perylene
GC/MS	EPA 625; EPA 8270C/D	Benzo(k)fluoranthene
GC/MS	EPA 8270C/D	Benzoic Acid
GC/MS	EPA 8270C/D	Benzyl alcohol
GC/MS	EPA 8270C/D	1,1-Biphenyl
GC/MS	EPA 625; EPA 8270C/D	bis(2-Chloroethoxy)methane
GC/MS	EPA 625; EPA 8270C/D	bis(2-Chloroethyl) ether
GC/MS	EPA 625; EPA 8270C/D	bis(2-Chloroisopropyl) ether (2, 2`-Oxybis(1-chloropropane)
GC/MS	EPA 625; EPA 8270C/D	bis(2-Ethylhexyl)adipate
GC/MS	EPA 625; EPA 8270C/D	bis(2-Ethylhexyl) phthalate (DEHP)
GC/MS	EPA 625; EPA 8270C/D	Butyl benzyl phthalate
GC/MS	EPA 8270C/D	Caprolactam
GC/MS	EPA 8270C/D	Carbazole
GC/MS	EPA 8270C/D	Chlorobenzilate
GC/MS	EPA 625; EPA 8270C/D	Chrysene
GC/MS	EPA 8270C/D	Diallate
GC/MS	EPA 8270C/D	Dibenzo(a,j)acridine
GC/MS	EPA 625; EPA 8270C/D	Dibenz(a h)anthracene
GC/MS	EPA 8270C/D	Dibenzofuran
GC/MS	EPA 8270C/D	Diethyladipate
GC/MS	EPA 625; EPA 8270C/D	Diethyl phthalate

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-Potable Water		
Technology	Method	Analyte
GC/MS	EPA 8270C/D	Dimethoate
GC/MS	EPA 625; EPA 8270C/D	Dimethyl phthalate
GC/MS	EPA 625; EPA 8270C/D	Di-n-butyl phthalate
GC/MS	EPA 625; EPA 8270C/D	Di-n-octyl phthalate
GC/MS	EPA 8270C/D	Dinoseb
GC/MS	EPA 8270C/D	Disulfoton
GC/MS	EPA 8270C/D	Ethyl methanesulfonate
GC/MS	EPA 8270C/D	Ethyl parathion
GC/MS	EPA 8270C/D	Ethyl methacrylate
GC/MS	EPA 8270C/D	Famfur
GC/MS	EPA 625; EPA 8270C/D	Fluoranthene
GC/MS	EPA 625; EPA 8270C/D	Fluorene
GC/MS	EPA 625; EPA 8270C/D	Hexachlorobenzene
GC/MS	EPA 625; EPA 8270C/D	Hexachlorobutadiene
GC/MS	EPA 625; EPA 8270C/D	Hexachlorocyclopentadiene
GC/MS	EPA 625; EPA 8270C/D	Hexachloroethane
GC/MS	EPA 8270C/D	Hexachlorophene
GC/MS	EPA 8270C/D	Hexachloropropene
GC/MS	EPA 625; EPA 8270C/D	Indeno(1, 2, 3-cd)pyrene
GC/MS	EPA 8270C/D	Isodrin
GC/MS	EPA 625; EPA 8270C/D	Isophorone
GC/MS	EPA 8270C/D	Isosafrole
GC/MS	EPA 8270C/D	Kepone
GC/MS	EPA 8270C/D	Methapyriline
GC/MS	EPA 8270C/D	Methy methanesulfonate
GC/MS	EPA 8270C/D	Methyl parathion
GC/MS	EPA 625; EPA 8270C/D	Naphthalene
GC/MS	EPA 625; EPA 8270C/D	Nitrobenzene
GC/MS	EPA 8270C/D	Nitroquinoline-1-oxide
GC/MS	EPA 8270C/D	n-Nitrosodiethylamine
GC/MS	EPA 625; EPA 8270C/D	n-Nitrosodimethylamine

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-Potable Water		
Гесhnology	Method	Analyte
GC/MS	EPA 8270C/D	n-Nitroso-di-n-butylamine
GC/MS	EPA 625; EPA 8270C/D	n-Nitrosodi-n-propylamine
GC/MS	EPA 625; EPA 8270C/D	n-Nitrosodiphenylamine
GC/MS	EPA 8270C/D	n-Nitrosomethylethylamine
GC/MS	EPA 8270C/D	n-Nitrosomorpholine
GC/MS	EPA 8270C/D	n-Nitrosopiperidine
GC/MS	EPA 8270C/D	n-Nitrosopyrrolidine
GC/MS	EPA 8270C/D	O,O,O-Triethyl phosphorothioate
GC/MS	EPA 8270C/D	o,o-Diethyl o-2pyrazinyl phosphorothioate
GC/MS	EPA 8270C/D	o-Toluidine
GC/MS	EPA 8270C/D	Pentachlorobenzene
GC/MS	EPA 8270C/D	Pentachloronitrobenzene
GC/MS	EPA 625; EPA 8270C/D	Pentachlorophenol
GC/MS	EPA 8270C/D	Phenacetin
GC/MS	EPA 625; EPA 8270C/D	Phenanthrene
GC/MS	EPA 625; EPA 8270C/D	Phenol
GC/MS	EPA 8270C/D	Phorate
GC/MS	EPA 8270C/D	Pronamide
GC/MS	EPA 625; EPA 8270C/D	Pyrene
GC/MS	EPA 8270C/D	Pyridine
GC/MS	EPA 8270C/D	Safrole
GC/MS	EPA 8270C/D	Sulfotepp
GC/MS	EPA 8270C/D	Thionazin
GC/MS	EPA 625; EPA 8270C/D	3, 3'-Dichlorobenzidine
GC/MS	EPA 8270C/D	3, 3'-Dimethylbenzidine
GC/MS	EPA 8270C/D SIM	1,1'-Biphenyl
GC/MS	EPA 8270C/D SIM	1,2,4,5-Tetrachlorobenzene
GC/MS	EPA 8270C/D SIM	1,4-Dioxane
GC/MS	EPA 8270C/D SIM	1-Methylnaphthalene
GC/MS	EPA 8270C/D SIM	2,2'-Oxybis(1-chloropropane
GC/MS	EPA 8270C/D SIM	2,3,4,6-Tetrachlorophenol

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n-Potable Water		
Technology	Method	Analyte
GC/MS	EPA 8270C/D SIM	2,4,5-Trichlorophenol
GC/MS	EPA 8270C/D SIM	2,4,6-Trichlorophenol
GC/MS	EPA 8270C/D SIM	2,4-Dichlorophenol
GC/MS	EPA 8270C/D SIM	2,4-Dimethylphenol
GC/MS	EPA 8270C/D SIM	2,4-Dinitrophenol
GC/MS	EPA 8270C/D SIM	2,4-Dinitrotoluene
GC/MS	EPA 8270C/D SIM	2,6-Dinitrotoluene
GC/MS	EPA 8270C/D SIM	2-Chloronaphthalene
GC/MS	EPA 8270C/D SIM	2-Chlorophenol
GC/MS	EPA 8270C/D SIM	2-Methylnaphthalene
GC/MS	EPA 8270C/D SIM	2-Methylphenol
GC/MS	EPA 8270C/D SIM	2-Nitroaniline
GC/MS	EPA 8270C/D SIM	2-Nitrophenol
GC/MS	EPA 8270C/D SIM	3&4-Methylphenol
GC/MS	EPA 8270C/D SIM	3,3'-Dichlorobenzidine
GC/MS	EPA 8270C/D SIM	3-Nitroaniline
GC/MS	EPA 8270C/D SIM	4,6-Dinitro-2-methylphenol
GC/MS	EPA 8270C/D SIM	4-Bromophenyl-phenylether
GC/MS	EPA 8270C/D SIM	4-Chloro-3-methylphenol
GC/MS	EPA 8270C/D SIM	4-Chloroaniline
GC/MS	EPA 8270C/D SIM	4-Chlorophenyl-phenylether
GC/MS	EPA 8270C/D SIM	4-Nitroaniline
GC/MS	EPA 8270C/D SIM	4-Nitrophenol
GC/MS	EPA 8270C/D SIM	Acenaphthene
GC/MS	EPA 8270C/D SIM	Acenaphthylene
GC/MS	EPA 8270C/D SIM	Acetophenone
GC/MS	EPA 8270C/D SIM	Anthracene
GC/MS	EPA 8270C/D SIM	Atrazine
GC/MS	EPA 8270C/D SIM	Benzaldehyde
GC/MS	EPA 8270C/D SIM	Benzo(a)anthracene
GC/MS	EPA 8270C/D SIM	Benzo(a)pyrene

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Toolando	M-41 3	1 14
Technology	Method	Analyte
GC/MS	EPA 8270C/D SIM	Benzo(b)fluoranthene
GC/MS	EPA 8270C/D SIM	Benzo(g,h,i)perylene
GC/MS	EPA 8270C/D SIM	Benzo(k)fluoranthene
GC/MS	EPA 8270C/D SIM	Bis(2-chloroethoxy)methane
GC/MS	EPA 8270C/D SIM	Bis(2-chloroethyl)ether
GC/MS	EPA 8270C/D SIM	Bis(2-ethylhexyl)phthalate
GC/MS	EPA 8270C/D SIM	Butylbenzylphthalate
GC/MS	EPA 8270C/D SIM	Caprolactam
GC/MS	EPA 8270C/D SIM	Carbazole
GC/MS	EPA 8270C/D SIM	Chrysene
GC/MS	EPA 8270C/D SIM	Dibenzo(a,h)anthracene
GC/MS	EPA 8270C/D SIM	Dibenzofuran
GC/MS	EPA 8270C/D SIM	Diethylphthalate
GC/MS	EPA 8270C/D SIM	Dimethyl phthalate
GC/MS	EPA 8270C/D SIM	Di-n-butylphthalate
GC/MS	EPA 8270C/D SIM	Di-n-octylphthalate
GC/MS	EPA 8270C/D SIM	Fluoranthene
GC/MS	EPA 8270C/D SIM	Fluorene
GC/MS	EPA 8270C/D SIM	Hexachlorobenzene
GC/MS	EPA 8270C/D SIM	Hexachlorobutadiene
GC/MS	EPA 8270C/D SIM	Hexachlorocyclopentadiene
GC/MS	EPA 8270C/D SIM	Hexachloroethane
GC/MS	EPA 8270C/D SIM	Indeno(1,2,3-cd)pyrene
GC/MS	EPA 8270C/D SIM	Isophorone
GC/MS	EPA 8270C/D SIM	Naphthalene
GC/MS	EPA 8270C/D SIM	Nitrobenzene
GC/MS	EPA 8270C/D SIM	n-Nitroso-di-n-propylamine
GC/MS	EPA 8270C/D SIM	n-Nitrosodiphenylamine
GC/MS	EPA 8270C/D SIM	Pentachlorophenol
GC/MS	EPA 8270C/D SIM	Phenanthrene
GC/MS	EPA 8270C/D SIM	Phenol

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Non-Potable Water		
Technology	Method	Analyte
GC/MS	EPA 8270C/D SIM	Pyrene
HPLC/UV	EPA 8330A/B	1, 3, 5-Trinitrobenzene
HPLC/UV	EPA 8330A/B	1, 3-Dinitrobenzene
HPLC/UV	EPA 8330A/B	2, 4, 6-Trinitrotoluene
HPLC/UV	EPA 8330A/B	2, 4-Dinitrotoluene
HPLC/UV	EPA 8330A/B	2, 6-Dinitrotoluene
HPLC/UV	EPA 8330A/B	2-Amino-4, 6 -Dinitrotoluene
HPLC/UV	EPA 8330A/B	2-Nitrotoluene
HPLC/UV	EPA 8330A/B	3-Nitrotoluene
HPLC/UV	EPA 8330A/B	3,5-Dinitroaniline
HPLC/UV	EPA 8330A/B	4-Amino-2, 6-Dinitrotoluene
HPLC/UV	EPA 8330A/B	4-Nitrotoluene
HPLC/UV	EPA 8330A/B	Ethylene glycol dinitrate (EGDN)
HPLC/UV	EPA 8330A/B	Hexahydro-1, 3, 5-trinitro-1, 3, 5-triazine (RDX)
HPLC/UV	EPA 8330A/B	Nitrobenzene
HPLC/UV	EPA 8330A MOD	Nitroglycerin
HPLC/UV	EPA 8330B	Nitroglycerin
HPLC/UV	EPA 8330A/B	Octahydro-1, 3, 5, 7-tetrazocine (HMX)
HPLC/UV	EPA 8330A/B	Pentaerythritol Tetranitrate (PETN)
HPLC/UV	EPA 8330A/B	Tetryl
CVAA	EPA 245.1; EPA 7470A	Mercury
CVAF	EPA 1631E	Low Level Mercury
ICP/AES	EPA 200.7; EPA 6010B/C	Aluminum
ICP/AES	EPA 200.7; EPA 6010B/C	Antimony
ICP/AES	EPA 200.7; EPA 6010B/C	Arsenic
ICP/AES	EPA 200.7; EPA 6010B/C	Barium
ICP/AES	EPA 200.7; EPA 6010B/C	Beryllium
ICP/AES	EPA 200.7; EPA 6010B/C	Boron
ICP/AES	EPA 200.7; EPA 6010B/C	Cadmium
ICP/AES	EPA 200.7; EPA 6010B/C	Calcium
ICP/AES	EPA 200.7; EPA 6010B/C	Chromium

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Non-Potable Water		
Technology	Method	Analyte
ICP/AES	EPA 200.7; EPA 6010B/C	Cobalt
ICP/AES	EPA 200.7; EPA 6010B/C	Copper
ICP/AES	EPA 200.7; EPA 6010B/C	Iron
ICP/AES	EPA 200.7; EPA 6010B/C	Lead
ICP/AES	EPA 200.7; EPA 6010B/C	Magnesium
ICP/AES	EPA 200.7; EPA 6010B/C	Manganese
ICP/AES	EPA 200.7; EPA 6010B/C	Molybdenum
ICP/AES	EPA 200.7; EPA 6010B/C	Nickel
ICP/AES	EPA 200.7; EPA 6010B/C	Potassium
ICP/AES	EPA 200.7; EPA 6010B/C	Selenium
ICP/AES	EPA 200.7; EPA 6010B/C	Silicon
ICP/AES	EPA 200.7; EPA 6010B/C	Silver
ICP/AES	EPA 200.7; EPA 6010B/C	Sodium
ICP/AES	EPA 6010B/C	Strontium
ICP/AES	EPA 200.7; EPA 6010B/C	Thallium
ICP/AES	EPA 200.7; EPA 6010B/C	Tin
ICP/AES	EPA 200.7; EPA 6010B/C	Titanium
ICP/AES	EPA 200.7; EPA 6010B/C	Vanadium
ICP/AES	EPA 200.7; EPA 6010B/C	Zinc
ICP/MS	EPA 200.8; EPA 6020A	Aluminum
ICP/MS	EPA 200.8; EPA 6020A	Antimony
ICP/MS	EPA 200.8; EPA 6020A	Arsenic
ICP/MS	EPA 200.8; EPA 6020A	Barium
ICP/MS	EPA 200.8; EPA 6020A	Beryllium
ICP/MS	EPA 200.8; EPA 6020A	Boron
ICP/MS	EPA 200.8; EPA 6020A	Cadmium
ICP/MS	EPA 200.8; EPA 6020A	Calcium
ICP/MS	EPA 200.8; EPA 6020A	Chromium
ICP/MS	EPA 200.8; EPA 6020A	Cobalt
ICP/MS	EPA 200.8; EPA 6020A	Copper
ICP/MS	EPA 200.8; EPA 6020A	Iron

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Non-Potable Water		
Technology	Method	Analyte
ICP/MS	EPA 200.8; EPA 6020A	Lead
ICP/MS	EPA 200.8; EPA 6020A	Magnesium
ICP/MS	EPA 200.8; EPA 6020A	Manganese
ICP/MS	EPA 200.8; EPA 6020A	Molybdenum
ICP/MS	EPA 200.8; EPA 6020A	Nickel
ICP/MS	EPA 200.8; EPA 6020A	Potassium
ICP/MS	EPA 200.8; EPA 6020A	Selenium
ICP/MS	EPA 200.8; EPA 6020A	Silicon
ICP/MS	EPA 200.8; EPA 6020A	Silver
ICP/MS	EPA 200.8; EPA 6020A	Sodium
ICP/MS	EPA 6020A	Strontium
ICP/MS	EPA 200.8; EPA 6020A	Thallium
ICP/MS	EPA 200.8; EPA 6020A	Tin
ICP/MS	EPA 200.8; EPA 6020A	Titanium
ICP/MS	EPA 200.8; EPA 6020A	Tungsten
ICP/MS	EPA 200.8	Uranium
ICP/MS	EPA 200.8; EPA 6020A	Vanadium
ICP/MS	EPA 200.8; EPA 6020A	Zinc
IC	EPA 300.0; EPA 9056A	Bromide
IC	EPA 300.0; EPA 9056A	Chloride
IC	EPA 300.0; EPA 9056A	Fluoride
IC	EPA 300.0; EPA 9056A	Nitrate as N
IC	EPA 300.0; EPA 9056A	Nitrite as N
IC	EPA 300.0; EPA 9056A	Nitrate + Nitrite
IC	EPA 300.0; EPA 9056A	Orthophosphate as P
IC	EPA 300.0; EPA 9056A	Sulfate
IC	SOP CA-776	Lactic Acid
IC	SOP CA-776	Acetic Acid
IC	SOP CA-776	Propionic Acid
IC	SOP CA-776	Formic Acid
IC	SOP CA-776	Butyric Acid

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on-Potable Water		
Technology	Method	Analyte
IC	SOP CA-776	Pyruvic Acid
IC	SOP CA-776	i-Pentanoic Acid
IC	SOP CA-776	Pentanoic Acid
IC	SOP CA-776	i-Hexanoic Acid
IC	SOP CA-776	Hexanoic Acid
Titration	EPA 310.1; SM 2320B	Alkalinity
Caculation	SM 2340B	Hardness
Gravimetric	EPA 1664A; EPA 9070A	Oil and Grease, Oil and Grease with SGT
Gravimetric	SM 2540B/C/D	Solids
ISE	EPA 120.1; SM 2510B	Conductivity
ISE	SM 2520B	Practical Salinity
ISE	SM 4500F- C	Fluoride
ISE	SM 4500H+ B	рН
ISE	SM 5210B	TBOD / CBOD
Physical	EPA 1010A	Ignitability
Physical	EPA 9040C	pH
Titration	SM 2340C	Hardness
Titration	SM 4500SO <sub>3</sub> B	Sulfite
Titration	EPA 9034; SM 4500S <sup>2-</sup> F	Sulfide
Titration	EPA SW-846 Chapter 7.3.4	Reactive Sulfide
IR	EPA 9060A; SM 5310B	Total organic carbon
Turbidimetric	EPA 180.1; SM 2130B	Turbidity
Turbidimetric	EPA 9038; ASTM 516-02	Sulfate
UV/VIS	EPA 335.4; EPA 9012B; SM 4500-CN G	Amenable cyanide
UV/VIS	EPA 350.1; SM 4500NH3 H	Ammonia as N
UV/VIS	SM 3500Fe D	Ferrous Iron
UV/VIS	EPA 351.2	Kjeldahl nitrogen - total
UV/VIS	EPA 353.2; SM 4500NO3 F	Nitrate + Nitrite
UV/VIS	EPA 353.2; SM 4500NO3 F	Nitrate as N
UV/VIS	EPA 353.2; SM 4500NO3 F	Nitrite as N
UV/VIS	EPA 365.2; SM 4500P E	Orthophosphate as P

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Non-Potable Water		
Technology	Method	Analyte
UV/VIS	EPA 365.4	Phosphorus total
UV/VIS	EPA 821/R-91-100	AVS-SEM
UV/VIS	EPA 410.4	COD
UV/VIS	EPA 420.1; EPA 9065	Total Phenolics
UV/VIS	SM 4500Cl G	Total Residual Chlorine
UV/VIS	SM 5540C	MBAS
UV/VIS	EPA 7196A; SM 3500-Cr D	Chromium VI
UV/VIS	EPA 9012B; EPA 335.4	Total Cyanide
UV/VIS	EPA 9251; SM 4500Cl E	Chloride
UV/VIS	EPA SW-846 Chapter 7.3.4	Reactive Cyanide

Preparation	Method	Туре
Cleanup Methods	EPA 3640A	Gel Permeation Clean-up
Cleanup Methods	EPA 3630C	Silica Gel
Cleanup Methods	EPA 3660B	Sulfur Clean-Up
Cleanup Methods	EPA 3665A	Sulfuric Acid Clean-Up
Organic Preparation	EPA 3510C	Separatory Funnel Extraction
Organic Preparation	EPA 3520C	Continuous Liquid-Liquid Extraction
Inorganic Preparation	EPA 3010A	Hotblock
Volatile Organic Preparation	EPA 5030C	Purge and Trap

Solid and Chemical Waste			
Technology	Method	Analyte	
GC/ECD	EPA 8081B	2,4`-DDD	
GC/ECD	EPA 8081B	2,4`-DDE	
GC/ECD	EPA 8081B	2,4`-DDT	
GC/ECD	EPA 8081B	4, 4`-DDD	
GC/ECD	EPA 8081B	4, 4`-DDE	
GC/ECD	EPA 8081B	4, 4`-DDT	
GC/ECD	EPA 8081B	Aldrin	

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Гесhnology	Method	Analyte
GC/ECD	EPA 8081B	alpha-BHC (alpha-Hexachlorocyclohexane)
GC/ECD	EPA 8081B	Alpha-Chlordane
GC/ECD	EPA 8081B	beta-BHC (beta-Hexachlorocyclohexane)
GC/ECD	EPA 608; EPA 8081B	Chlordane (tech.)
GC/ECD	EPA 8081B	Cis-Nonachlor
GC/ECD	EPA 8081B	delta-BHC
GC/ECD	EPA 8081B	Dieldrin
GC/ECD	EPA 8081B	Endosulfan I
GC/ECD	EPA 8081B	Endosulfan II
GC/ECD	EPA 8081B	Endosulfan sulfate
GC/ECD	EPA 8081B	Endrin
GC/ECD	EPA 8081B	Endrin aldehyde
GC/ECD	EPA 8081B	Endrin Ketone
GC/ECD	EPA 8081B	gamma-BHC (Lindane gamma- Hexachlorocyclohexane)
GC/ECD	EPA 8081B	gamma-Chlordane
GC/ECD	EPA 8081B	Heptachlor
GC/ECD	EPA 8081B	Heptachlor epoxide
GC/ECD	EPA 8081B	Hexachlorobenzene
GC/ECD	EPA 8081B	Methoxychlor
GC/ECD	EPA 8081B	Mirex
GC/ECD	EPA 8081B	Oxychlordane
GC/ECD	EPA 8081B	Toxaphene (Chlorinated camphene)
GC/ECD	EPA 8081B	Trans-Nonachlor
GC/ECD	EPA 8082A	Aroclor-1016 (PCB-1016)
GC/ECD	EPA 8082A	Aroclor-1221 (PCB-1221)
GC/ECD	EPA 8082A	Aroclor-1232 (PCB-1232)
GC/ECD	EPA 8082A	Aroclor-1242 (PCB-1242)
GC/ECD	EPA 8082A	Aroclor-1248 (PCB-1248)
GC/ECD	EPA 8082A	Aroclor-1254 (PCB-1254)
GC/ECD	EPA 8082A	Aroclor-1260 (PCB-1260)

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id and Chemical Waste		
Technology	Method	Analyte
GC/ECD	EPA 8082A MOD	Aroclor-1268 (PCB-1268)
GC/ECD	EPA 8082A	2, 2', 3, 3', 4, 4', 5, 5', 6-Nonachlorobiphenyl (BZ 206)
GC/ECD	EPA 8082A	2, 2', 3, 3', 4, 4', 5, 6-Octachlorobiphenyl (BZ 19
GC/ECD	EPA 8082A	2, 2', 3, 3', 4, 4', 5-Heptachlorobiphenyl (BZ 170
GC/ECD	EPA 8082A	2, 2', 3, 3', 4, 4'-Hexachlorobiphenyl (BZ 128)
GC/ECD	EPA 8082A	2, 2', 3, 4, 4', 5, 5'-Heptachlorobiphenyl (BZ 180
GC/ECD	EPA 8082A	2, 2', 3, 4, 4', 5', 6-Heptachlorobiphenyl (BZ 183
GC/ECD	EPA 8082A	2, 2', 3, 4, 4', 5-Hexachlorobiphenyl (BZ 138)
GC/ECD	EPA 8082A	2, 2', 3, 4, 4', 6, 6'-Heptachlorobiphenyl (BZ 184
GC/ECD	EPA 8082A	2, 2', 3, 4', 5, 5', 6-Heptachlorobiphenyl (BZ 187
GC/ECD	EPA 8082A	2, 2', 3, 4, 5'-Pentachlorobiphenyl (BZ 87)
GC/ECD	EPA 8082A	2, 2', 3, 5'-Tetrachlorobiphenyl (BZ 44)
GC/ECD	EPA 8082A	2, 2', 4, 4', 5, 5'-Hexachlorobiphenyl (BZ 153)
GC/ECD	EPA 8082A	2, 2', 4, 5, 5'-Pentachlorobiphenyl (BZ 101)
GC/ECD	EPA 8082A	2, 2', 4, 5-Tetrachlorobiphenyl (BZ 48)
GC/ECD	EPA 8082A	2, 2', 4, 5'-Tetrachlorobiphenyl (BZ 49)
GC/ECD	EPA 8082A	2, 2', 5, 5'-Tetrachlorobiphenyl (BZ 52)
GC/ECD	EPA 8082A	2, 2', 5-Trichlorobiphenyl (BZ 18)
GC/ECD	EPA 8082A	2, 3, 3', 4, 4', 5-Hexachlorobiphenyl (BZ 156)
GC/ECD	EPA 8082A	2, 3, 3', 4, 4', 5'-Hexachlorobiphenyl (BZ 157)
GC/ECD	EPA 8082A	2, 3, 3', 4, 4'-Pentachlorobiphenyl (BZ 105)
GC/ECD	EPA 8082A	2, 3, 3', 4, 4', 5, 5'-Heptachlorobiphenyl (BZ 18
GC/ECD	EPA 8082A	2, 3', 4, 4', 5, 5'-Hexachlorobiphenyl (BZ 167)
GC/ECD	EPA 8082A	2, 3', 4, 4', 5-Pentachlorobiphenyl (BZ 118)
GC/ECD	EPA 8082A	2, 3', 4, 4',5'-Pentachlorobiphenyl (BZ 123)
GC/ECD	EPA 8082A	2, 3', 4, 4'-Tetrachlorobiphenyl (BZ 66)
GC/ECD	EPA 8082A	2, 3, 4, 4', 5-Pentachlorobiphenyl (BZ 114)
GC/ECD	EPA 8082A	2, 4, 4'-Trichlorobiphenyl (BZ 28)
GC/ECD	EPA 8082A	2, 4'-Dichlorobiphenyl (BZ 8)
GC/ECD	EPA 8082A	3, 3', 4, 4', 5, 5'-Hexachlorobiphenyl (BZ 169)
GC/ECD	EPA 8082A	3, 3', 4, 4', 5-Pentachlorobiphenyl (BZ 126)

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olid and Chemical Waste		
Technology	Method	Analyte
GC/ECD	EPA 8082A	3, 3', 4, 4'-Tetrachlorobiphenyl (BZ 77)
GC/ECD	EPA 8082A	3, 4, 4', 5-Tetrachlorobiphenyl (BZ 81)
GC/ECD	EPA 8082A	Decachlorobiphenyl (BZ 209)
GC/ECD	EPA 8151A	2, 4, 5-T
GC/ECD	EPA 8151A	2, 4-D
GC/ECD	EPA 8151A	2, 4-DB
GC/ECD	EPA 8151A	Dalapon
GC/ECD	EPA 8151A	Dicamba
GC/ECD	EPA 8151A	Dichloroprop
GC/ECD	EPA 8151A	Dinoseb
GC/ECD	EPA 8151A	MCPA
GC/ECD	EPA 8151A	МСРР
GC/ECD	EPA 8151A	Pentachlorophenol
GC/ECD	EPA 8151A	Silvex (2, 4, 5-TP)
GC/FID	EPA 8015C/D	Diesel range organics (DRO)
GC/FID	EPA 8015C/D	Total Petroleum Hydrocarbons (TPH)
GC/FID	EPA 8015C/D	Gasoline range organics (GRO)
GC/FID/PID	MA DEP VPH	Volatile Organic Hydrocarbons
GC/FID	MA DEP EPH	Extractable Petroleum Hydrocarbons
GC/FID	MA DEP EPH EPA 3546	Extractable Petroleum Hydrocarbons Microwave Extraction Preparation
GC/FID	СТ-ЕТРН	Total Petroleum Hydrocarbons
GC/FID	TNRCC Method 1005	Total Petroleum Hydrocarbons
GC/FID	FL-PRO	Petroleum Range Organics
GC/ECD	EPA 8011	1, 2-Dibromoethane (EDB)
GC/ECD	EPA 8011	1, 2-Dibromo-3-chloropropane
GC/MS	EPA 8260B/C	1, 1, 1, 2-Tetrachloroethane
GC/MS	EPA 8260B/C	1,1,2-Trichloro-1,2,2-trifluoroethane
GC/MS	EPA 8260B/C	1, 1, 1-Trichloroethane
GC/MS	EPA 8260B/C	1, 1, 2, 2-Tetrachloroethane
GC/MS	EPA 8260B/C	1, 1, 2-Trichloroethane
GC/MS	EPA 8260B/C	1, 1-Dichloroethane

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Solid and Chemical Waste	lid and Chemical Waste		
Technology	Method	Analyte	
GC/MS	EPA 8260B/C	1, 1-Dichloroethylene	
GC/MS	EPA 8260B/C	1, 1-Dichloropropene	
GC/MS	EPA 8260B/C	1, 2, 3-Trichlorobenzene	
GC/MS	EPA 8260B/C	1, 2, 3-Trichloropropane	
GC/MS	EPA 8260B/C	1,2,3-Trimethylbenzene	
GC/MS	EPA 8260B/C	1, 2, 4-Trichlorobenzene	
GC/MS	EPA 8260B/C	1, 2, 4-Trimethylbenzene	
GC/MS	EPA 8260B/C	1, 2-Dibromo-3-chloropropane	
GC/MS	EPA 8260B/C	1, 2-Dibromoethane	
GC/MS	EPA 8260B/C	1, 2-Dichlorobenzene	
GC/MS	EPA 8260B/C	1, 2-Dichloroethane	
GC/MS	EPA 8260B/C	1, 2-Dichloropropane	
GC/MS	EPA 8260B/C	1,3,5-Trichlorobenzene	
GC/MS	EPA 8260B/C	1, 3, 5-Trimethylbenzene	
GC/MS	EPA 8260B/C	1, 3-Dichlorobenzene	
GC/MS	EPA 8260B/C	1, 3-Dichloropropane	
GC/MS	EPA 8260B/C	1, 4-Dichlorobenzene	
GC/MS	EPA 8260B/C	1, 4-Dioxane	
GC/MS	EPA 8260B/C	1-Chlorohexane	
GC/MS	EPA 8260B/C	2, 2-Dichloropropane	
GC/MS	EPA 8260B/C	2-Butanone	
GC/MS	EPA 8260B/C	2-Chloroethyl vinyl ether	
GC/MS	EPA 8260B/C	2-Chlorotoluene	
GC/MS	EPA 8260B/C	2-Hexanone	
GC/MS	EPA 8260B/C	4-Chlorotoluene	
GC/MS	EPA 8260B/C	4-Methyl-2-pentanone	
GC/MS	EPA 8260B/C	Acetone	
GC/MS	EPA 8260B/C	Acetonitrile	
GC/MS	EPA 8260B/C	Acrolein	
GC/MS	EPA 8260B/C	Acrylonitrile	
GC/MS	EPA 8260B/C	Allyl chloride	

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Гесhnology	Method	Analyte
GC/MS	EPA 8260B/C	Benzene
GC/MS	EPA 8260B/C	Benzyl chloride
GC/MS	EPA 8260B/C	Bromobenzene
GC/MS	EPA 8260B/C	Bromochloromethane
GC/MS	EPA 8260B/C	Bromodichloromethane
GC/MS	EPA 8260B/C	Bromoform
GC/MS	EPA 8260B/C	Carbon disulfide
GC/MS	EPA 8260B/C	Carbon tetrachloride
GC/MS	EPA 8260B/C	Chlorobenzene
GC/MS	EPA 8260B/C	Chloroethane
GC/MS	EPA 8260B/C	Chloroform
GC/MS	EPA 8260B/C	Chloroprene
GC/MS	EPA 8260B/C	cis-1, 2-Dichloroethene
GC/MS	EPA 8260B/C	cis-1, 3-Dichloropropene
GC/MS	EPA 8260B/C	cis-1,3-Dichloro-2-butene
GC/MS	EPA 8260B/C	Cyclohexane
GC/MS	EPA 8260B/C	Dibromochloromethane
GC/MS	EPA 8260B/C	Dibromomethane
GC/MS	EPA 8260B/C	Dichlorodifluoromethane
GC/MS	EPA 8260B/C	Diethyl ether
GC/MS	EPA 8260B/C	Di-isopropylether
GC/MS	EPA 8260B/C	1,2-Dibromoethane (EDB)
GC/MS	EPA 8260B/C	Ethyl methacrylate
GC/MS	EPA 8260B/C	Ethylbenzene
GC/MS	EPA 8260B/C	Ethyl-t-butylether
GC/MS	EPA 8260B/C	Hexachlorobutadiene
GC/MS	EPA 8260B/C	Iodomethane
GC/MS	EPA 8260B/C	Isobutyl alcohol
GC/MS	EPA 8260B/C	Isopropyl alcohol
GC/MS	EPA 8260B/C	Isopropyl benzene

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Cechnology	Method	Analyte
GC/MS	EPA 8260B, C	Methyl acetate
GC/MS	EPA 8260B/C	Methacrylonitrile
GC/MS	EPA 8260B/C	Methyl bromide (Bromomethane)
GC/MS	EPA 8260B/C	Methyl chloride (Chloromethane)
GC/MS	EPA 8260B/C	Methyl methacrylate
GC/MS	EPA 8260B/C	Methyl tert-butyl ether
GC/MS	EPA 8260B/C	Methylcyclohexane
GC/MS	EPA 8260B/C	Methylene chloride
GC/MS	EPA 8260B/C	Naphthalene
GC/MS	EPA 8260B/C	n-Butylbenzene
GC/MS	EPA 8260B/C	n-proplybenzene
GC/MS	EPA 8260B/C	o-Xylene
GC/MS	EPA 8260B/C	pentachloroethane
GC/MS	EPA 8260B/C	p-Isopropyltoluene
GC/MS	EPA 8260B/C	Propionitrile
GC/MS	EPA 8260B/C	sec-butylbenzene
GC/MS	EPA 8260B/C	Styrene
GC/MS	EPA 8260B/C	t-Amylmethylether
GC/MS	EPA 8260B/C	tert-Butyl alcohol
GC/MS	EPA 8260B/C	tert-Butylbenzene
GC/MS	EPA 8260B/C	Tetrachloroethylene (Perchloroethylene)
GC/MS	EPA 8260B/C	Tetrahydrofuran
GC/MS	EPA 8260B/C	Toluene
GC/MS	EPA 8260B/C	trans-1, 2-Dichloroethylene
GC/MS	EPA 8260B/C	trans-1, 3-Dichloropropylene
GC/MS	EPA 8260B/C	Trans-1, 4-Dichloro-2-butuene
GC/MS	EPA 8260B/C	Trichloroethene (Trichloroethylene)
GC/MS	EPA 8260B/C	Trichlorofluoromethane
GC/MS	EPA 8260B/C	Vinyl acetate
GC/MS	EPA 8260B/C	Vinyl chloride
GC/MS	EPA 8260B/C	Xylene

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echnology	Method	Analyte
GC/MS	EPA 8260B/C SIM	1,1,1,2-Tetrachloroethane
GC/MS	EPA 8260B/C SIM	1,1,1-Trichloroethane
GC/MS	EPA 8260B/C SIM	1,1,2,2-Tetrachloroethane
GC/MS	EPA 8260B/C SIM	1, 1, 2-Trichloroethane
GC/MS	EPA 8260B/C SIM	1,2,3-Trichloropropane
GC/MS	EPA 8260B/C SIM	1,1-Dichloroethane
GC/MS	EPA 8260B/C SIM	1,1-Dichloroethene
GC/MS	EPA 8260B/C SIM	1,2,4-Trichlorobenzene
GC/MS	EPA 8260B/C SIM	1,2,4-Trimethylbenzene
GC/MS	EPA 8260B/C SIM	1,2-Dibromo-3-chloropropane
GC/MS	EPA 8260B/C SIM	1,2-Dibromoethane
GC/MS	EPA 8260B/C SIM	1,2-Dichlorobenzene
GC/MS	EPA 8260B/C SIM	1,2-Dichloroethane
GC/MS	EPA 8260B/C SIM	1,2-Dichloropropane
GC/MS	EPA 8260B/C SIM	1,3-Dichlorobenzene
GC/MS	EPA 8260B/C SIM	1,3-Dichloropropane
GC/MS	EPA 8260B/C SIM	1,4-Dichlorobenzene
GC/MS	EPA 8260B/C SIM	2-Hexanone
GC/MS	EPA 8260B/C SIM	4-Methyl-2-pentanone
GC/MS	EPA 8260B/C SIM	Benzene
GC/MS	EPA 8260B/C SIM	Bromodichloromethane
GC/MS	EPA 8260B/C SIM	Carbon Tetrachloride
GC/MS	EPA 8260B/C SIM	Chloroform
GC/MS	EPA 8260B/C SIM	Chloromethane
GC/MS	EPA 8260B/C SIM	cis-1,2-Dichloroethene
GC/MS	EPA 8260B/C SIM	cis-1,3-Dichloropropene
GC/MS	EPA 8260B/C SIM	Dibromochloromethane
GC/MS	EPA 8260B/C SIM	Ethylbenzene
GC/MS	EPA 8260B/C SIM	Isopropylbenzene
GC/MS	EPA 8260B/C SIM	Hexachlorobutadiene
GC/MS	EPA 8260B/C SIM	Methylcyclohexane

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Гесhnology	Method	Analyte
GC/MS	EPA 8260B/C SIM	m,p-Xylene
GC/MS	EPA 8260B/C SIM	o-Xylene
GC/MS	EPA 8260B/C SIM	Tetrachloroethene
GC/MS	EPA 8260B/C SIM	trans-1,2-Dichloroethene
GC/MS	EPA 8260B/C SIM	Trans-1,3-Dichloropropene
GC/MS	EPA 8260B/C SIM	Trichloroethene
GC/MS	EPA 8260B/C SIM	Trichlorofluoromethane
GC/MS	EPA 8260B/C SIM	Vinyl Chloride
GC/MS	EPA 8260B/C SIM	Xylenes (total)
GC/MS	EPA 8270C/D	1, 2, 4, 5-Tetrachlorobenzene
GC/MS	EPA 8270C/D	1, 2, 4-Trichlorobenzene
GC/MS	EPA 8270C/D	1, 2-Dichlorobenzene
GC/MS	EPA 8270C/D	1, 2-Diphenylhydrazine
GC/MS	EPA 8270C/D	1, 3, 5-Trinitrobenzene
GC/MS	EPA 8270C/D	1, 3-Dichlorobenzene
GC/MS	EPA 8270C/D	1, 3-Dinitrobenzene
GC/MS	EPA 8270C/D	1, 4-Dichlorobenzene
GC/MS	EPA 8270C/D	1, 4-Dioxane
GC/MS	EPA 8270C/D	1, 4-Naphthoquinone
GC/MS	EPA 8270C/D	1, 4-Phenylenediamine
GC/MS	EPA 8270C/D	1,1-Biphenyl
GC/MS	EPA 8270C/D	1-Chloronaphthalene
GC/MS	EPA 8270C/D	1-Methylnaphthalene
GC/MS	EPA 8270C/D	1-Naphthylamine
GC/MS	EPA 8270C/D	2, 3, 4, 6-Tetrachlorophenol
GC/MS	EPA 8270C/D	2, 4, 5-Trochlorophenol
GC/MS	EPA 8270C/D	2, 4, 6-Trichlorophenol
GC/MS	EPA 8270C/D	2, 4-Dichlorophenol
GC/MS	EPA 8270C/D	2, 4-Dimethylphenol
GC/MS	EPA 8270C/D	2, 4-Dinitrophenol

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Гесhnology	Method	Analyte
GC/MS	EPA 8270C/D	2, 6-Dichlorophenol
GC/MS	EPA 8270C/D	2, 6-Dinitrotoluene (2 6-DNT)
GC/MS	EPA 8270C/D	2-Acetylaminofluorene
GC/MS	EPA 8270C/D	2-Chloronaphthalene
GC/MS	EPA 8270C/D	2-Chlorophenol
GC/MS	EPA 8270C/D	2-Methyl-4, 6-dinitrophenol
GC/MS	EPA 8270C/D	2-Methylnaphthalene
GC/MS	EPA 8270C/D	2-Methylphenol
GC/MS	EPA 8270C/D	2-Naphthylamine
GC/MS	EPA 8270C/D	2-Nitroaniline
GC/MS	EPA 8270C/D	2-Nitrophenol
GC/MS	EPA 8270C/D	2-Picoline
GC/MS	EPA 8270C/D	3, 3`-Dichlorobenzidine
GC/MS	EPA 8270C/D	3, 3'-Dimethylbenzidine
GC/MS	EPA 8270C/D	3,4-Methylphenol
GC/MS	EPA 8270C/D	3-Methylcholanthrene
GC/MS	EPA 8270C/D	3-Nitroaniline
GC/MS	EPA 8270C/D	4-Aminobiphenyl
GC/MS	EPA 8270C/D	4-Bromophenyl phenyl ether
GC/MS	EPA 8270C/D	4-Chloro-3-methylphenol
GC/MS	EPA 8270C/D	4-Chloroaniline
GC/MS	EPA 8270C/D	4-Chlorophenyl phenylether
GC/MS	EPA 8270C/D	4-Dimethyl aminoazobenzene
GC/MS	EPA 8270C/D	4-Nitroaniline
GC/MS	EPA 8270C/D	4-Nitrophenol
GC/MS	EPA 8270C/D	4-Nitroquinoline-1-oxide
GC/MS	EPA 8270C/D	5-Nitro-o-toluidine
GC/MS	EPA 8270C/D	7,12-Dimethylbenz(a)anthracene
GC/MS	EPA 8270C/D	a a-Dimethylphenethylamine
GC/MS	EPA 8270C/D	Acenaphthene

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Technology	Method	Analyte
GC/MS	EPA 8270C/D	Acetophenone
GC/MS	EPA 8270C/D	Aniline
GC/MS	EPA 8270C/D	Anthracene
GC/MS	EPA 8270C/D	Aramite
GC/MS	EPA 8270C/D	Atrazine
GC/MS	EPA 8270C/D	Azobenzene
GC/MS	EPA 8270C/D	Benzaldehyde
GC/MS	EPA 8270C/D	Benzidine
GC/MS	EPA 8270C/D	Benzo(a)anthracene
GC/MS	EPA 8270C/D	Benzo(a)pyrene
GC/MS	EPA 8270C/D	Benzo(b)fluoranthene
GC/MS	EPA 8270C/D	Benzo(g h i)perylene
GC/MS	EPA 8270C/D	Benzo(k)fluoranthene
GC/MS	EPA 8270C/D	Benzoic Acid
GC/MS	EPA 8270C/D	Benzyl alcohol
GC/MS	EPA 8270C/D	bis(2-Chloroethoxy)methane
GC/MS	EPA 8270C/D	bis(2-Chloroethyl) ether
GC/MS	EPA 8270C/D	bis(2-Chloroisopropyl) ether (2, 2`-Oxybis(1-chloropropane))
GC/MS	EPA 8270C/D	bis(2-Ethylhexyl) phthalate (DEHP)
GC/MS	EPA 625; EPA 8270C/D	Bis(2-Ethylhexyl)adipate
GC/MS	EPA 8270C/D	Butyl benzyl phthalate
GC/MS	EPA 8270C/D	Caprolactam
GC/MS	EPA 8270C/D	Carbazole
GC/MS	EPA 8270C/D	Chlorobenzilate
GC/MS	EPA 8270C/D	Chrysene
GC/MS	EPA 8270C/D	Diallate
GC/MS	EPA 8270C/D	Dibenz(a h)anthracene
GC/MS	EPA 8270C/D	Dibenzo(a,j)acridine
GC/MS	EPA 8270C/D	Dibenzofuran
GC/MS	EPA 8270C/D	Diethyl phthalate
GC/MS	EPA 8270C/D	Diethyladipate

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lid and Chemical Waste		
Technology	Method	Analyte
GC/MS	EPA 8270C/D	Dimethoate
GC/MS	EPA 8270C/D	Dimethyl phthalate
GC/MS	EPA 8270C/D	Di-n-butyl phthalate
GC/MS	EPA 8270C/D	Di-n-octyl phthalate
GC/MS	EPA 8270C/D	Dinoseb
GC/MS	EPA 8270C/D	Disulfoton
GC/MS	EPA 8270C/D	Ethyl methacrylate
GC/MS	EPA 8270C/D	Ethyl methanesulfonate
GC/MS	EPA 8270C/D	Ethyl parathion
GC/MS	EPA 8270C/D	Famfur
GC/MS	EPA 8270C/D	Fluoranthene
GC/MS	EPA 8270C/D	Fluorene
GC/MS	EPA 8270C/D	Hexachlorobenzene
GC/MS	EPA 8270C/D	Hexachlorobutadiene
GC/MS	EPA 8270C/D	Hexachlorocyclopentadiene
GC/MS	EPA 8270C/D	Hexachloroethane
GC/MS	EPA 8270C/D	Hexachlorophene
GC/MS	EPA 8270C/D	Hexachloropropene
GC/MS	EPA 8270C/D	Indeno(1, 2, 3-cd)pyrene
GC/MS	EPA 8270C/D	Isodrin
GC/MS	EPA 8270C/D	Isophorone
GC/MS	EPA 8270C/D	Isosafrole
GC/MS	EPA 8270C/D	Kepone
GC/MS	EPA 8270C/D	Methapyriline
GC/MS	EPA 8270C/D	Methyl methanesulfonate
GC/MS	EPA 8270C/D	Methyl parathion
GC/MS	EPA 8270C/D	Naphthalene
GC/MS	EPA 8270C/D	Nitrobenzene
GC/MS	EPA 8270C/D	n-Nitrosodiethylamine
GC/MS	EPA 8270C/D	n-Nitrosodimethylamine
GC/MS	EPA 8270C/D	n-Nitroso-di-n-butylamine

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Гесhnology	Method	Analyte
GC/MS	EPA 8270C/D	n-Nitrosodi-n-propylamine
GC/MS	EPA 8270C/D	n-Nitrosodiphenylamine
GC/MS	EPA 8270C/D	n-Nitrosomethylethylamine
GC/MS	EPA 8270C/D	n-Nitrosomorpholine
GC/MS	EPA 8270C/D	n-Nitrosopiperidine
GC/MS	EPA 8270C/D	n-Nitrosopyrrolidine
GC/MS	EPA 8270C/D	O, O, O-Triethyl phosphorothioate
GC/MS	EPA 8270C/D	o,o-Diethyl o-2-pyrazinyl phosphorothioate
GC/MS	EPA 8270C/D	o-Toluidine
GC/MS	EPA 8270C/D	Pentachlorobenzene
GC/MS	EPA 8270C/D	Pentachloronitrobenzene
GC/MS	EPA 8270C/D	Pentachlorophenol
GC/MS	EPA 8270C/D	Phenacetin
GC/MS	EPA 8270C/D	Phenanthrene
GC/MS	EPA 8270C/D	Phenol
GC/MS	EPA 8270C/D	Phorate
GC/MS	EPA 8270C/D	Pronamide
GC/MS	EPA 8270C/D	Pyrene
GC/MS	EPA 8270C/D	Pyridine
GC/MS	EPA 8270C/D	Safrole
GC/MS	EPA 8270C/D	Sulfotepp
GC/MS	EPA 8270C/D	Thionazin
GC/MS	EPA 8270C/D SIM	1,1'-Biphenyl
GC/MS	EPA 8270C/D SIM	1,2,4,5-Tetrachlorobenzene
GC/MS	EPA 8270C/D SIM	1,4-Dioxane
GC/MS	EPA 8270C/D SIM	1-Methylnaphthalene
GC/MS	EPA 8270C/D SIM	2,2'-Oxybis(1-chloropropane
GC/MS	EPA 8270C/D SIM	2,3,4,6-Tetrachlorophenol
GC/MS	EPA 8270C/D SIM	2,4,5-Trichlorophenol
GC/MS	EPA 8270C/D SIM	2,4,6-Trichlorophenol

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d and Chemical Waste			
Technology	Method	Analyte	
GC/MS	EPA 8270C/D SIM	2,4-Dimethylphenol	
GC/MS	EPA 8270C/D SIM	2,4-Dinitrophenol	
GC/MS	EPA 8270C/D SIM	2,4-Dinitrotoluene	
GC/MS	EPA 8270C/D SIM	2,6-Dinitrotoluene	
GC/MS	EPA 8270C/D SIM	2-Chloronaphthalene	
GC/MS	EPA 8270C/D SIM	2-Chlorophenol	
GC/MS	EPA 8270C/D SIM	2-Methylnaphthalene	
GC/MS	EPA 8270C/D SIM	2-Methylphenol	
GC/MS	EPA 8270C/D SIM	2-Nitroaniline	
GC/MS	EPA 8270C/D SIM	2-Nitrophenol	
GC/MS	EPA 8270C/D SIM	3&4-Methylphenol	
GC/MS	EPA 8270C/D SIM	3,3'-Dichlorobenzidine	
GC/MS	EPA 8270C/D SIM	3-Nitroaniline	
GC/MS	EPA 8270C/D SIM	4,6-Dinitro-2-methylphenol	
GC/MS	EPA 8270C/D SIM	4-Bromophenyl-phenylether	
GC/MS	EPA 8270C/D SIM	4-Chloro-3-methylphenol	
GC/MS	EPA 8270C/D SIM	4-Chloroaniline	
GC/MS	EPA 8270C/D SIM	4-Chlorophenyl-phenylether	
GC/MS	EPA 8270C/D SIM	4-Nitroaniline	
GC/MS	EPA 8270C/D SIM	4-Nitrophenol	
GC/MS	EPA 8270C/D SIM	Acenaphthene	
GC/MS	EPA 8270C/D SIM	Acenaphthylene	
GC/MS	EPA 8270C/D SIM	Acetophenone	
GC/MS	EPA 8270C/D SIM	Anthracene	
GC/MS	EPA 8270C/D SIM	Atrazine	
GC/MS	EPA 8270C/D SIM	Benzaldehyde	
GC/MS	EPA 8270C/D SIM	Benzo(a)anthracene	
GC/MS	EPA 8270C/D SIM	Benzo(a)pyrene	
GC/MS	EPA 8270C/D SIM	Benzo(b)fluoranthene	
GC/MS	EPA 8270C/D SIM	Benzo(g,h,i)perylene	
GC/MS	EPA 8270C/D SIM	Benzo(k)fluoranthene	

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Гесhnology	Method	Analyte
GC/MS	EPA 8270C/D SIM	Bis(2-chloroethoxy)methane
GC/MS	EPA 8270C/D SIM	Bis(2-chloroethyl)ether
GC/MS	EPA 8270C/D SIM	Bis(2-ethylhexyl)phthalate
GC/MS	EPA 8270C/D SIM	Butylbenzylphthalate
GC/MS	EPA 8270C/D SIM	Caprolactam
GC/MS	EPA 8270C/D SIM	Carbazole
GC/MS	EPA 8270C/D SIM	Chrysene
GC/MS	EPA 8270C/D SIM	Dibenzo(a,h)anthracene
GC/MS	EPA 8270C/D SIM	Dibenzofuran
GC/MS	EPA 8270C/D SIM	Diethylphthalate
GC/MS	EPA 8270C/D SIM	Dimethyl phthalate
GC/MS	EPA 8270C/D SIM	Di-n-butylphthalate
GC/MS	EPA 8270C/D SIM	Di-n-octylphthalate
GC/MS	EPA 8270C/D SIM	Fluoranthene
GC/MS	EPA 8270C/D SIM	Fluorene
GC/MS	EPA 8270C/D SIM	Hexachlorobenzene
GC/MS	EPA 8270C/D SIM	Hexachlorobutadiene
GC/MS	EPA 8270C/D SIM	Hexachlorocyclopentadiene
GC/MS	EPA 8270C/D SIM	Hexachloroethane
GC/MS	EPA 8270C/D SIM	Indeno(1,2,3-cd)pyrene
GC/MS	EPA 8270C/D SIM	Isophorone
GC/MS	EPA 8270C/D SIM	Naphthalene
GC/MS	EPA 8270C/D SIM	Nitrobenzene
GC/MS	EPA 8270C/D SIM	n-Nitroso-di-n-propylamine
GC/MS	EPA 8270C/D SIM	n-Nitrosodiphenylamine
GC/MS	EPA 8270C/D SIM	Pentachlorophenol
GC/MS	EPA 8270C/D SIM	Phenanthrene
GC/MS	EPA 8270C/D SIM	Phenol
GC/MS	EPA 8270C/D SIM	Pyrene
HPLC/UV	EPA 8330A	1 ,3, 5-Trinitrobenzene
HPLC/UV	EPA 8330A	1, 3-Dinitrobenzene

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olid and Chemical W	aste		
Technology	Method	Analyte	
HPLC/UV	EPA 8330A	2, 4, 6-Trinitrotoluene	
HPLC/UV	EPA 8330A	2, 4-Dinitrotoluene	
HPLC/UV	EPA 8330A	2, 6-Dinitrotoluene	
HPLC/UV	EPA 8330A	2-Amino-4, 6-dinitrotoluene	
HPLC/UV	EPA 8330A	2-Nitrotoluene	
HPLC/UV	EPA 8330A	3-Nitrotoluene	
HPLC/UV	EPA 8330A	3,5-Dinitroaniline	
HPLC/UV	EPA 8330A	4-Amino-2,6-dinitrotoluene	
HPLC/UV	EPA 8330A	4-Nitrotoluene	
HPLC/UV	EPA 8330A	Ethylene glycol dinitrate (EGDN)	
HPLC/UV	EPA 8330A	Hexahydr-1, 3, 5-trinitro-1, 3, 5-triazine (RDX)	
HPLC/UV	EPA 8330A	Nitrobenzene	
HPLC/UV	EPA 8330A MOD	Nitroglycerin	
HPLC/UV	EPA 8330A	Octahydro-1, 3, 5, 7-tetrazocine (HMX)	
HPLC/UV	EPA 8330A	Pentaerythritol Tetranitrate (PETN)	
HPLC/UV	EPA 8330A	Tetryl	
HPLC/UV	8330B (W/O Soil Grinding)	1, 3, 5-Trinitrobenzene	
HPLC/UV	8330B (W/O Soil Grinding)	1, 3-Dinitrobenzene	
HPLC/UV	8330B (W/O Soil Grinding)	2, 4, 6-Trinitrotoluene	
HPLC/UV	8330B (W/O Soil Grinding)	2, 4-Dinitrotoluene	
HPLC/UV	8330B (W/O Soil Grinding)	2, 6-Dinitrotoluene	
HPLC/UV	8330B (W/O Soil Grinding)	2-Amino-4, 6 –Dinitrotoluene	
HPLC/UV	8330B (W/O Soil Grinding)	2-Nitrotoluene	
HPLC/UV	8330B (W/O Soil Grinding)	3-Nitrotoluene	
HPLC/UV	8330B (W/O Soil Grinding)	3,5-Dinitroaniline	
HPLC/UV	8330B (W/O Soil Grinding)	4-Amino-2,3-Dinitrotoluene	
HPLC/UV	8330B (W/O Soil Grinding)	4-Nitrotoluene	
HPLC/UV	8330B (W/O Soil Grinding)	Ethylene glycol dinitrate (EGDN)	
HPLC/UV	8330B (W/O Soil Grinding)	Hexahydr-1, 3, 5-trinitro-1, 3, 5-triazine (RDX)	
HPLC/UV	8330B (W/O Soil Grinding)	Nitrobenzene	
HPLC/UV	8330B (W/O Soil Grinding)	Nitroglycerin	

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Solid and Chemical Waste		
Technology	Method	Analyte
HPLC/UV	8330B (W/O Soil Grinding)	Octahydro-1, 3, 5, 7-tetrazocine (HMX)
HPLC/UV	8330B (W/O Soil Grinding)	Pentaerythritol Tetranitrate (PETN)
HPLC/UV	8330B (W/O Soil Grinding)	Tetryl
CVAA	EPA 7471B	Mercury
CVAF	EPA 1631E	Low Level Mercury
ICP/AES	EPA 6010B/C	Aluminum
ICP/AES	EPA 6010B/C	Antimony
ICP/AES	EPA 6010B/C	Arsenic
ICP/AES	EPA 6010B/C	Barium
ICP/AES	EPA 6010B/C	Beryllium
ICP/AES	EPA 6010B/C	Boron
ICP/AES	EPA 6010B/C	Cadmium
ICP/AES	EPA 6010B/C	Calcium
ICP/AES	EPA 6010B/C	Chromium
ICP/AES	EPA 6010B/C	Cobalt
ICP/AES	EPA 6010B/C	Copper
ICP/AES	EPA 6010B/C	Iron
ICP/AES	EPA 6010B/C	Lead
ICP/AES	EPA 6010B/C	Magnesium
ICP/AES	EPA 6010B/C	Manganese
ICP/AES	EPA 6010B/C	Molybdenum
ICP/AES	EPA 6010B/C	Nickel
ICP/AES	EPA 6010B/C	Potassium
ICP/AES	EPA 6010B/C	Selenium
ICP/AES	EPA 6010B/C	Silicon
ICP/AES	EPA 6010B/C	Silver
ICP/AES	EPA 6010B/C	Sodium
ICP/AES	EPA 6010B/C	Strontium
ICP/AES	EPA 6010B/C	Thallium
ICP/AES	EPA 6010B/C	Tin
ICP/AES	EPA 6010B/C	Titanium

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olid and Chemical Waste		
Technology	Method	Analyte
ICP/AES	EPA 6010B/C	Vanadium
ICP/AES	EPA 6010B/C	Zinc
ICP/MS	EPA 6020A	Aluminum
ICP/MS	EPA 6020A	Antimony
ICP/MS	EPA 6020A	Arsenic
ICP/MS	EPA 6020A	Barium
ICP/MS	EPA 6020A	Beryllium
ICP/MS	EPA 6020A	Boron
ICP/MS	EPA 6020A	Cadmium
ICP/MS	EPA 6020A	Calcium
ICP/MS	EPA 6020A	Chromium
ICP/MS	EPA 6020A	Cobalt
ICP/MS	EPA 6020A	Copper
ICP/MS	EPA 6020A	Iron
ICP/MS	EPA 6020A	Lead
ICP/MS	EPA 6020A	Magnesium
ICP/MS	EPA 6020A	Manganese
ICP/MS	EPA 6020A	Molybdenum
ICP/MS	EPA 6020A	Nickel
ICP/MS	EPA 6020A	Potassium
ICP/MS	EPA 6020A	Selenium
ICP/MS	EPA 6020A	Silver
ICP/MS	EPA 6020A	Sodium
ICP/MS	EPA 6020A	Strontium
ICP/MS	EPA 6020A	Thallium
ICP/MS	EPA 6020A	Tin
ICP/MS	EPA 6020A	Titanium
ICP/MS	EPA 6020A	Tungsten
ICP/MS	EPA 6020A	Vanadium
ICP/MS	EPA 6020A	Zinc
IC	EPA 9056A	Chloride

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Technology	Method	Analyte	
IC	EPA 9056A	Fluoride	
IC	EPA 9056A	Nitrate as N	
IC	EPA 9056A	Nitrite as N	
IC	EPA 9056A	Orthophosphate	
IC	EPA 9056A	Sulfate	
Gravimetric	EPA 9071A; EPA 9071B	Oil and Grease, Oil and Grease with SGT	
Physical	EPA 1010A	Ignitability	
Physical	EPA 9045D	pH	
Titration	EPA SW-846 Chapter 7.3.4	Reactive Sulfide	
Titration	Walkley-Black	Total Organic Carbon	
IR	Lloyd Kahn	Total organic carbon	
Turbidimetric	EPA 9038; ASTM 516-02	Sulfate	
UV/VIS	EPA 350.1; SM 4500NH3 H	Ammonia as N	
UV/VIS	EPA 9251; SM 4500Cl E	Chloride	
UV/VIS	EPA SW-846 Chapter 7.3.4	Reactive Cyanide	
UV/VIS	EPA 821/R-91-100	AVS-SEM	
UV/VIS	SM 3500Fe D	Ferrous Iron	
Cleanup Methods	EPA 3630C	Silica Gel	
UV/VIS	EPA 7196	Chromium VI	
UV/VIS	EPA 7196A	Chromium VI	
UV/VIS	EPA 9012B	Total cyanide	
Preparation	Method	Туре	
Preparation	EPA 1311	Toxicity Characteristic Leaching Procedure	
Preparation	EPA 1312	Synthetic Precipitation Leaching Procedure	
Cleanup Methods	EPA 3660B	Sulfur Clean-up	
Cleanup Methods	EPA 3620C	Florsil Clean-up	
Cleanup Methods	EPA 3630C	Silica Gel Clean-up	
Cleanup Methods	EPA 3640A	GPC Clean-up	
Organic Preparation	EPA 3540C	Soxhlet Extraction	
Organic Preparation	EPA 3545A	Pressurized Fluid Extraction	

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Solid and Chemical Waste				
Technology	Method	Analyte		
Organic Preparation	EPA 3546	Microwave Extraction Preparation for EPA 8082A, 8081B and 8270C, D		
Organic Preparation	EPA 3550C	Sonication		
Inorganics Preparation	EPA 3050B	Hotblock		
Inorganics Preparation	EPA 3060A	Alkaline Digestion		
Volatile Organics Preparation	EPA 5035/5035A	Closed System Purge and Trap		

Air			
Technology	Method	Analyte	
GC/MS	EPA TO-15	Propene	
GC/MS	EPA TO-15	1, 1, 1-Trichloroethane	
GC/MS	EPA TO-15	1, 1, 2, 2-Tetrachloroethane	
GC/MS	EPA TO-15	1, 1, 2-Trichloroethane	
GC/MS	EPA TO-15	1, 1-Dichloroethane	
GC/MS	EPA TO-15	1, 1-Dichloroethylene	
GC/MS	EPA TO-15	1, 2, 4-Trichlorobenzene	
GC/MS	EPA TO-15	1, 2, 4-Trimethylbenzene	
GC/MS	EPA TO-15	1, 2-Dibromoethane (EDB)	
GC/MS	EPA TO-15	1,2-Dichloro-1,1,2,2-tetrafluoroethane (Freon 114)	
GC/MS	EPA TO-15	1, 2-Dichlorobenzene	
GC/MS	EPA TO-15	1, 2-Dichloroethane	
GC/MS	EPA TO-15	1, 2-Dichloroethenes (Total)	
GC/MS	EPA TO-15	1, 2-Dichloropropane	
GC/MS	EPA TO-15	1, 3, 5-Trimethylbenzene	
GC/MS	EPA TO-15	1, 3-Butadiene	
GC/MS	EPA TO-15	1, 3-Dichlorobenzene	
GC/MS	EPA TO-15	1, 4-Dichlorobenzene	
GC/MS	EPA TO-15	1,4-Difluorobenzene	
GC/MS	EPA TO-15	1, 4-Dioxane	
GC/MS	EPA TO-15	2-Butanone	
GC/MS	EPA TO-15	2-Hexanone	

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Technology	Method	Analyte	
GC/MS	EPA TO-15	2-Propanol	
GC/MS	EPA TO-15	4-Ethyltoluene	
GC/MS	EPA TO-15	4-Methyl-2-pentanone	
GC/MS	EPA TO-15	Acetone	
GC/MS	EPA TO-15	Acrolein	
GC/MS	EPA TO-15	Benzene	
GC/MS	EPA TO-15	Benzyl chloride	
GC/MS	EPA TO-15	Bromochloromethane	
GC/MS	EPA TO-15	Bromodichloromethane	
GC/MS	EPA TO-15	Bromoform	
GC/MS	EPA TO-15	Carbon disulfide	
GC/MS	EPA TO-15	Carbon tetrachloride	
GC/MS	EPA TO-15	Chlorobenzene	
GC/MS	EPA TO-15	Chloroethane	
GC/MS	EPA TO-15	Chloroform	
GC/MS	EPA TO-15	Cis-1, 2-Dichloroethene	
GC/MS	EPA TO-15	Cis-1, 3-Dichloropropene	
GC/MS	EPA TO-15	Cyclohexane	
GC/MS	EPA TO-15	Dibromochloromethane	
GC/MS	EPA TO-15	Dichlorodifluoromethane (Freon 12)	
GC/MS	EPA TO-15	Ethanol	
GC/MS	EPA TO-15	Ethyl acetate	
GC/MS	EPA TO-15	Ethylbenzene	
GC/MS	EPA TO-15	Hexachlorobutadiene	
GC/MS	EPA TO-15	Isopropyl alcohol	
GC/MS	EPA TO-15	m, p-Xylene	
GC/MS	EPA TO-15	Methyl bromide (Bromomethane)	
GC/MS	EPA TO-15	Methyl chloride (Chloromethane)	
GC/MS	EPA TO-15	Methyl methacrylate	
GC/MS	EPA TO-15	Methyl tert-butyl ether	
GC/MS	EPA TO-15	Methylene chloride	

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Air		
Technology	Method	Analyte
GC/MS	EPA TO-15	Naphthalene
GC/MS	EPA TO-15	n-Heptane
GC/MS	EPA TO-15	n-Hexane
GC/MS	EPA TO-15	o-Xylene
GC/MS	EPA TO-15	Styrene
GC/MS	EPA TO-15	Tetrachloroethylene (Perchloroethylene)
GC/MS	EPA TO-15	Tetrahydrofuran
GC/MS	EPA TO-15	Toluene
GC/MS	EPA TO-15	trans-1, 2-Dichloroethylene
GC/MS	EPA TO-15	trans-1, 3-Dichloropropylene
GC/MS	EPA TO-15	Trichloroethene (Trichloroethylene)
GC/MS	EPA TO-15	Trichlorofluoromethane (Freon 11)
GC/MS	EPA TO-15	1,1,2-Trichloro1,2,2-trifluoroethane (Freon 113)
GC/MS	EPA TO-15	Vinyl acetate
GC/MS	EPA TO-15	Vinyl chloride
GC/MS	EPA TO-15	Xylenes (Total)

#### Notes:

1) This laboratory offers commercial testing service.

Approved by:

R. Douglas Leonard
Chief Technical Officer

Re-issued: 2/1/13

Date: February 1, 2013

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# State of Rhode Island and Providence Plantations DEPARTMENT OF HEALTH Certifies

## KATAHDIN ANALYTICAL SERVICES INC 600 TECHNOLOGY WAY SCARBOROUGH ME 04074 Laboratory Director: DEBORAH NADEAU

for the analysis of:

Potable Water Organic Chemistry - Potable Water Inorganic Chemistry - Non-potable Water Organic

Chemistry - Non-potable Water Inorganic Chemistry -

This certificate is issued, pursuant to Rhode Island General Laws 23-16.2 and supersedes all previous Rhode Island certificates issued to this laboratory. Certification is no guarantee of the validity of the laboratory results.

This certificate is valid only when accompanied by the certificate and list of analytes and methods for which certification has been granted based upon the following out of state certification(s):

Certifying Authority
MAINE
NH

Certification Number ME0019 200112 Expiration Date 06/01/2014 04/02/2013

KATAHDIN ANALYTICAL SERVICES INC is responsible for maintaining each of the certifications listed above. Failure to notify the Laboratory Certification Officer of any change in the status of these certifications may result in the suspension or revocation of certification. Contact the Laboratory Certification Officer to verify the current certification status of this laboratory.

Michael Fine, MD Director of Health

Expires: 12/30/2013

# KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-520 Revision History Cover Page Page 1

	ATION OF AQUEOUS SAMPLES FOR ANALEUM HYDROCARBONS or DIESEL RANGE		
Prepared By:	Mike Thomas	Date:	6/97
Approved By:			
Group Supervisor:	michal F: Thomas	Date:_	1/29/01
Operations Manager	: Joh C. Benton	Date:_	1/29/01
QA Officer:	Detorah J. Nadeau	Date:_	1.29.01
General Manager:	Dunauf huskan	Date:_	1/29/01
	, 0		8
Revision History:			

COD			T .	
SOP	Changes	Approval	Approval	Effective
Revision		Initials	Date	Date
01	Format changes, added pollution prevention, removed 50:11 extraction, other minor changes to sections 7,8 and 04 Table.	9n	129,01	1/29/01
02	Wording added or changed to clarify sections 5,6,8,+9. Minor changes throughout. Hew figure.	HRC	11.08.04 H.08.04	11.08.04
03	Added - the condenser temperature during extraction.  Added - variable transformers output should be set at 55% updated Logbook	LAD	04/06	04/06
04	Added waste generated information. Added Acetone to reagents. Added QC pH must beadjusted. Added use of boiling stones in Sox. ext. Removed Maclz references. Updated Table 2. Added definitions	LAN	09107	09107
05	updated Logbook, added recording of lot numbers and nitrogen evaporation water both temperature in logbook	n LAP	11/08	1103

SOP Number: CA-520 Revision History Cover Page – Cont.

Page 2

TITLE: PREPARATION OF AQUEOUS SAMPLES FOR ANALYSIS OF EXTRACTABLE PETROLEUM HYDROCARBONS or DIESEL RANGE ORGANICS (DRO)

Revision History:

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
06	Added commercially available DRU product to section 5. updated logbook pagl.	LAD	08110	08/10
<b>6</b> 7	Minor changes to Section? to reflect Current practices. Removed fewled from filter papertype. Updated MOL, LOD, LOD information. Added Do Dand NECK Repeaced Removed Sect. 8.4 and added it to Method M.	LAO es,	04/12	04/12
			, , , , , , , , , , , , , , , , , , ,	

SOP Number: CA-520-07 Date Issued: 04/12

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#### 1.0 SCOPE AND APPLICATION

The purpose of this SOP is to describe the method used by Katahdin Analytical Services technical personnel to prepare samples for analyses of Total Petroleum Hydrocarbons (TPH) or Diesel Range Organics (DRO) in water. These compounds correspond to a hydrocarbon range of C9-C36 inclusive and a boiling point range between approximately 170°C and 430°C. The method is based on a solvent extraction procedure followed by Gas Chromatography (GC) analysis.

The method is designed to measure "mid-range" to "heavy" petroleum products. This range would include JP-4, JP-5, JP-8, kerosene, diesel #2, #4, #6, and motor oil. Components greater than C36 present in products are not readily amenable to this method. If, based on a review of the chromatogram, the presence of these product types is suspected, a qualitative description should be included in the report. Additional analyses may be performed including, but not limited to, analysis of additional reference materials. These additional efforts are not contained within this method.

#### 1.1 Definitions

TOTAL PETROLEUM HYDROCARBONS (TPH): All resolved and unresolved material eluting from n-nonane (n-C9) through n-hexatriacontane (n-C36), inclusive.

DIESEL RANGE ORGANICS (DRO): All resolved and unresolved material eluting from decane (n-C10) through n-octacosane (n-C28), inclusive.

ANALYTICAL BATCH: 20 or fewer samples which are analyzed together with the same method sequence and the same lots of reagents and with the manipulations common to each sample within the same time period or in continuous sequential time periods.

METHOD BLANK (laboratory reagent blank): An artificial sample designed to determine if method analytes or other interferences are present in the laboratory environment, the reagents, or the apparatus. For aqueous samples, reagent water is used as a blank matrix; for soil samples, baked organic-free sand is used as a blank matrix. The blank is taken through the appropriate steps of the process.

LABORATORY CONTROL SAMPLE (LCS): A blank that has been spiked with the analyte(s) from an independent source, and is analyzed exactly like a sample. Its purpose is to determine whether the methodology is in control, and whether the laboratory is capable of making accurate and precise measurements. The matrix used should be phase matched with the samples and well characterized.

MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD): Predetermined quantities of stock solutions of certain analytes are added to a sample matrix prior to sample extraction and analysis. Samples are split into duplicates, spiked and analyzed. Percent recoveries are calculated for each of the analytes detected. The relative

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percent difference between the samples is calculated and used to assess analytical precision.

SURROGATES: Organic compounds which are similar to analytes of interest in chemical composition, extraction and chromatography, but which are not normally found in environmental samples. These compounds are spiked into all blanks, standards, samples and spiked samples prior to analysis. Percent recoveries are calculated for each surrogate.

#### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of analysts experienced in the extraction of water samples for the determination of petroleum hydrocarbons or DROs. Each analyst must demonstrate and document their ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, current revision, "Personnel Training & Documentation of Capability," current revision, and Section 8.2.

It is the responsibility of all Katahdin technical personnel involved in the extraction of water and soil samples for the determination of petroleum hydrocarbons or DROs to read and understand this SOP, to adhere to the procedures outlined, and to properly document their data in the appropriate lab notebook. Any deviations from the test or irregularities with the samples should also be recorded in the lab notebook and reported to the Department Manager or designated qualified data reviewer responsible for this data.

It is the responsibility of the Department Manager to oversee that members of their group follow this SOP, to ensure that their work is properly documented and to initiate periodic review of the associated logbooks.

#### 1.3 Safety

Users of this procedure must be cognizant of inherent laboratory hazards, proper disposal procedures for contaminated materials and appropriate segregation of hazardous wastes. The toxicity or carcinogenicity of each reagent used in this method has not been precisely defined; however, each chemical should be treated as a potential health hazard. A reference file of material safety data sheets is available to all personnel involved in the chemical analysis. Everyone involved with the procedure must be familiar with the MSDSs for all the materials used in this procedure.

Each qualified analyst or technician must be familiar with Katahdin Analytical Environmental Health and Safety Manual including the Katahdin Hazardous Waste Plan and must follow appropriate procedures. These include the use of appropriate personal protective equipment (PPE) such as safety glasses, gloves and lab coats when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location of all safety equipment. Each analyst shall

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receive a safety orientation from their Department Manager, or designee, appropriate for the job functions they will perform.

#### 1.4 Pollution Prevention/Waste Disposal

Whenever possible, laboratory personnel should use pollution prevention techniques to address their waste generation. Refer to the current revision of the Katahdin Hazardous Waste Plan for further details on pollution prevention techniques.

Wastes generated during the preparation of samples must be disposed of in accordance with the Katahdin Hazardous Waste Plan and Safety Manual and SOP SD-903, "Sample Disposal," current revision. Expired standards are lab packed, placed in the Katahdin hazardous waste storage area, and disposed of in accordance with this SOP.

Any methylene chloride solvent waste generated during the rinsing of glassware etc. should be disposed of in the "D" waste stream satellite accumulation area nearest the point of generation. Acetone is considered flammable waste, and should be disposed of in the "O" waste stream satellite accumulation area nearest the point of generation. Sodium sulfate waste should be disposed of in the soil with organics "I" waste stream satellite accumulation area nearest the point of generation. Please refer to the current revision of SOP CA-107 for the location of satellite waste accumulation areas.

#### 2.0 SUMMARY OF METHOD

Samples are extracted with methylene chloride using separatory funnels following EPA Method 3510 or continuous liquid liquid extractors (CLLE) 3520 for subsequent analysis for Total Petroleum Hydrocarbons or Diesel Range Organics. The extract is dried, concentrated and injected into a capillary column gas chromatograph. This method is suitable for the analysis of aqueous samples.

This method is based in part on: 1) Petroleum Hydrocarbon Methods by API revised August 1993; 2) The Wisconsin DRO method, 3) SW-846 methods 3510, 3520, 3540, and 3550, 4) The Massachusetts EPH method; and 5) The Maine DRO method.

#### 3.0 INTERFERENCES

3.1 Other organic compounds including chlorinated hydrocarbons, phenols, and phthalate esters are measurable by this method. As defined in the method, the DRO results include these compounds. Interferences coextracted from the samples will vary considerably from source to source. If analysis of an extracted sample is prevented due to interferences, further cleanup of the sample extract may be needed to minimize interferences.

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3.2 Method interferences are reduced by washing all glassware with hot soapy water and then rinsing it sequentially with tap water, methanol or acetone, and methylene chloride. Method/reagent blanks (Surrogate Control Samples) must be analyzed with each batch to demonstrate that the samples are free from method interferences.

3.3 High purity reagents such as Burdick and Jackson GC methylene chloride or Baker capillary grade methylene chloride must be used to minimize interferences.

#### 4.0 APPARATUS AND MATERIALS

Prior to use, all glassware must be rinsed three times with the solvent to be used for extraction.

- 4.1 Separatory Funnel 2000 mL capacity, Nalgene Teflon FEP separatory funnels with Nalgene Tefzel® screw-cap closures (or equivalent)
- 4.2 Concentrator tube 10 mL, graduated
- 4.3 Evaporative flask Kuderna-Danish, 500 mL capacity attached to concentrator with neck clips
- 4.4 Snyder column Kuderna-Danish, three ball macro
- 4.5 Graduated cylinders 100 mL, 1000 mL, or 2000 mL
- 4.6 Short stem funnels
- 4.7 250 mL amber collection bottles with Teflon-lined caps
- 4.8 1.8 mL, 12 mL and/or 16 mL glass vials with Teflon-lined caps
- 4.9 Continuous liquid-liquid extractors (CLLE) including body, 500 mL flat bottom boiling flask and Alhin condensers
- 4.10 Filter paper, 18.5 cm, Fisher brand or Whatman #4 (or equivalent)
- 4.11 Nitrogen evaporation apparatus.
- 4.12 Boiling chips approximately 10/40 mesh, Teflon or selenized carborundum, 12 mesh (or equivalent).
- 4.13 Water bath eight position concentric ring bath or equivalent, equipped with a calibrated thermometer.

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#### 5.0 REAGENTS

- 5.1 Reagent water water in which an interferent is not observed at or above the PQL for any parameter of interest (laboratory reagent grade water or equivalent).
- 5.2 Sodium sulfate (granular, anhydrous and powdered, anhydrous) (ACS reagent grade), Na<sub>2</sub>SO<sub>4</sub>. Certified by the manufacturer/vendor as purified by heating at 400°C for 4 hours prior to receipt by the laboratory.
- 5.3 Acid for preserving water samples: A 1:1 mixture of reagent water and concentrated hydrochloric acid. Use ~5 mL per 1 L sample.
- 5.4 Methylene Chloride (MeCL<sub>2</sub>) and Acetone Pesticide grade or better. Lots must be verified by concentrating 300-400 mL to 1.0 mL and evaluating by GC/MS.
- 5.5 Surrogate spiking solution Prepare a solution of o-Terphenyl at a concentration of 20 ug/mL in acetone. Store the solution at -10 to -20 °C in a Teflon sealed container. Solution must be verified by GC/FID prior to use, and must be replaced every 6 months or sooner if degradation is evident.
- 5.6 Matrix Spike/Lab Control Sample spiking solution Prepare a matrix spiking solution in pesticide grade Acetone that contains all target analytes listed below:

Component	Concentration µg/mL
Decane	50
Dodecane	50
Tetradecane	50
Hexadecane	50
Octadecane	50
Eicosane	50
Docosane	50
Tetracosane	50
Hexacosane	50
Octacosane	50

**Alternatively,** a Matrix Spike/Lab Control Sample spiking solution may be prepared using a commercially available fuel oil such as Diesel Fuel #2, unweathered, from Restek. Prepare a 500 ug/L standard in pesticide grade Acetone. This spike may be required for by some certain states, federal programs, or clients (such as South Carolina).

Store the both solutions at -10 to -20 °C in a Teflon sealed container. The solutions must be verified by GC/ECD prior to use, and must be replaced every 6 months or sooner if degradation is evident.

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#### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

Aqueous samples are collected in a 1L amber glass bottle. As soon as possible after samples are received they must be stored at  $4^{\circ}$ C ( $\pm 2^{\circ}$ C) until extraction. When water samples are not received cold, the fact should be noted on the chain of custody form. The pH of aqueous samples must be checked with pH paper upon receipt to ensure that the samples have been acid preserved. If the pH is not < 2, the fact should be noted on the chain of custody form. Samples that have not been preserved should be preserved at this time and a notation made on the chain of custody form.

Holding time for extraction of aqueous samples for Methods 3510 and 3520 is 7 days from date of sample collection, although the analyst should be aware that actual holding times employed may be project/program specific.

Store all extracts at 4°C (±2°C) in labeled Teflon-sealed containers. See SOP SD-902, "Sample Receipt and Internal Control," current revision, for storage areas and temperature maintenance procedures.

#### 7.0 PROCEDURES

The following information must be recorded in the extraction logbook (all that are applicable).

- Extraction method
- Surrogate and spike IDs
- Lot numbers of all solvents, acids and bases, sodium sulfate, filter paper
- Nitrogen evaporation water bath temperature
- Ha •
- Extraction and concentration dates
- Extraction and concentration analyst
- Separatory funnel extraction start and end times.
- CLLE extraction start and end times, also the prep start and end times.
- Sample ID or QC sample ID
- Initial and final volumes
- Surrogate and spike amounts
- Final extract tray location
- Any comments regarding the sample extraction (ie. Emulsion)

#### SEPARATORY FUNNEL SAMPLE EXTRACTION

If an emulsion prevents acceptable recovery or client history indicates samples may demonstrate matrix interference, then samples should be extracted by continuous liquid-liquid extraction (CLLE).

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- 7.1 Rinse <u>all</u> glassware three times with methylene chloride prior to use.
- 7.2 Label a 2 L Teflon separatory funnel and a 250 mL amber collection bottle clearly. Label should include laboratory sample number, matrix, analyte, and extraction date. Be sure that the detachable stopcocks are secured to the separatory funnels before adding samples.
- 7.3 Measure the initial volume by comparing the meniscus of the sample with the reference bottle of the same bottle type. Please refer to SOP CA-108, "Basic Laboratory Technique", for the reference bottle verification procedure. Record the volume and any notable characteristics (e.g. color, presence of sediment, or odor) in the extraction logbook.
- 7.4 Transfer the contents of the sample bottle to a 2 L separatory funnel.
- 7.5 Transfer 1 L of reagent water to a 2 L separatory funnel. This serves as a method blank for the extraction batch. A method blank must be prepared for every daily extraction batch of twenty or fewer samples.
- 7.6 Transfer 1 L of reagent water to a 2 L separatory funnel. This will serve as a Laboratory Control Sample (LCS). An LCS is required for every daily extraction batch of twenty or fewer samples. If an MS/MSD pair is not extracted on a particular day, an LCS/LCSD pair may be required in order to meet client-specific or program-specific requirements. This information will be disseminated from the project manager or Department Manager.
- 7.7 A matrix spike/matrix spike duplicate (MS/MSD) is to be prepared as requested by a client or, at a minimum, one pair per 20 samples if there is sufficient sample volume. Measure the initial volume as in 7.3. Transfer two additional 1 L aliquots of sample to 2 L separatory funnels for a matrix spike and matrix spike duplicate (MS/MSD).
  - Note: Sufficient sample volume should be available without depleting all remaining sample aliquots.
- 7.8 Check to make sure the pH is <2. Note in the prep logbook if the pH is not <2 and adjust if needed with 1:1 HCl. (This should have been recorded and corrected, if necessary, at the time of sample receipt by the sample custodians.)
- 7.9 Adjust method blank and LCS/LCSD pH to <2 with 1:1 HCl.
- 7.10 Using a gas-tight syringe, add 1.0 mL of surrogate spiking solution to all samples the blank, LCS/LCSD and MS/MSD, if performed.
- 7.11 Using a gas-tight syringe, add 1.0 mL of matrix spiking solution to each LCS/LCSD and MS/MSD.

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7.12 Carefully add 60 mLs of methylene chloride to each empty sample bottle rinse the bottle and transfer the solvent into the appropriate separatory funnel making sure the dispenser tip does not come in contact with the bottle. Add 60 mL of methylene chloride directly to the blank and LCS/LCSD.

- 7.13 Ensure that each screw-cap is secured tightly to the separatory funnel to prevent leaks. Extract the sample by first shaking the funnel by hand for a few seconds, venting often in a hood to release pressure. Place funnel on mechanical shaker and shake for 3 minutes. Allow phases to separate. Drain the methylene chloride layer into the 250 mL amber collection bottle.
- 7.14 If an emulsion forms, mechanical techniques must be employed to achieve maximum separation and solvent recovery. Such means include swirling and centrifugation and draining through a small separatory funnel. In certain instances, transferring the entire sample into a continuous liquid-liquid extractor may be the only alternative. If any such techniques are used, they must be noted in the extractions logbook.
- 7.15 Add a second 60 mL aliquot of methylene chloride to the separatory funnel and extract for the second time (see 7.11 7.12). Collect the methylene chloride layer in the same 250 mL amber collection bottle.
- 7.16 Repeat the extraction for a third time as described in 7.13.
- 7.17 Proceed to Section 7.32 for extract concentration procedures.

#### CONTINUOUS LIQUID-LIQUID SAMPLE EXTRACTION (CLLE)

- 7.18 Set up the CLLE apparatus and add one or two boiling stones to the flask. All glassware should be rinsed three times with methylene chloride and the extract flasks properly labeled.
- 7.19 Add approximately 500 600 mL of methylene chloride to the CLLE body.
- 7.20 Add 1 L reagent water to a CLLE body. This is the method blank for this extraction batch. Be sure that no water leaks into the round bottom flask. A method blank is required for every extraction batch of twenty or fewer samples.
- 7.21 Prepare an LCS for every daily extraction batch of twenty or fewer samples. Add 1 L of reagent water to a CLLE body. If an MS/MSD pair is not extracted on a particular day, an LCS/LCSD pair may be required in order to meet client-specific or program-specific requirements. This information will be disseminated from the project manager or Department Manager.

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7.22 Check to make sure the pH is <2. Note in the prep logbook if the pH is not <2 and adjust if needed with 1:1 HCl. (This should have been recorded and corrected, if necessary, at the time of sample receipt by the sample custodians.)

- 7.23 Determine the initial volume as in 7.3. Transfer the samples to the CLLE bodies, being sure no sample leaks into the round bottom flask.
- 7.24 Transfer two 1 L portions of a sample to CLLE bodies for preparation of a matrix spike/matrix spike duplicate if required. An MS/MSD is required if requested by the client or per 20 samples or every 14 days, whichever occurs first. (Refer to the logbook page, "date QC expires"). Note: Sufficient sample volume should be available without depleting all remaining sample aliquots.
- 7.25 For each sample, rinse the original sample container with approximately 30 mL of methylene chloride being careful not to touch the bottle with the dispenser tip. Add this rinse to the CLLE body.
- 7.26 Adjust method blank and LCS/LCSD pH to <2 with 1:1 HCl.
- 7.27 Add 1.0 mL of the surrogate spiking solution to each sample including the blank, LCS/LCSD and MS/MSD, if performed.
- 7.29 Add 1.0 mL of matrix spiking solution to the appropriate LCS/LCSD and MS/MSD pair, if performed, and stir.
- 7.30 Rinse each 45/50 condenser joint with methylene chloride. Attach cooloing water Alhin condensers set to a temperature of 15 °C. Turn on the heating mantles, the rheostat of the variable transformers should be set to 55% of the output voltage. Allow the samples to extract for 20 ± 2 hours. Turn off the mantles and let samples cool.
- 7.31 Proceed to Section 7.32 for sample extract concentration procedures.

#### CONCENTRATION OF WATER SAMPLE EXTRACTS

- 7.32 Rinse the K-D glassware (flask, concentration tube, and Snyder column, including the ground glass joints on the flask and columns) three times with methylene chloride before assembling. Add a few boiling chips to the K-D. Insert 18.5 cm filter papers into short stem powder funnels and add ~ 2 inches of sodium sulfate crystals. Place the assembled K-D's under the funnels.
- 7.33 Transfer the methylene chloride extracts to the K-D concentrator setups through the sodium sulfate in the funnels. This is the drying step, which is required to remove residual water from the extracts. Any large water layers must be removed by other means, prior to pouring through the sodium sulfate. After pouring all of the extract

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volume through the sodium sulfate, rinse the extract bottle three times with  $\sim 2-3$  mLs of methylene chloride. Add the rinsings through the sodium sulfate to complete a quantitative transfer. Rinse the sodium sulfate with  $\sim 15$  mLs of methylene chloride and allow to drain.

- 7.34 Transfer the labels from the collection bottles or round bottom flasks (from the CLLE extraction) to the K-Ds. Remove the funnel and attach a 3-ball macro Snyder column. Pre-wet the Snyder column with 1 mL of methylene chloride.
- 7.35 Place the K-D in a hot water bath (75-85°C). Gently swirl K-D in the water until boiling begins. At the proper distillation rate, the Snyder balls should chatter but the chambers should not flood with condensed solvent. The K-D should be kept in a vertical orientation while on the bath. When the apparent volume in the concentrator tube reaches  $\approx$  5-6 mL, remove the K-D from the water bath. Allow the K-D to cool for 10 minutes. Rinse the Snyder column lower joint with  $\approx$  1 mL of methylene chloride. Remove the Snyder column. Wipe off any water from the neck above the lower joint of the flask. Separate the K-D flask from the concentrator tube, rinsing the ground glass joint with  $\approx$  1 mL methylene chloride.
- 7.36 Reduce the methylene chloride extract in the concentrator tube to approximately 1 mL using the nitrogen blow-down apparatus. The bath temperature must be no higher than the boiling point of the solvent (39°C for methylene chloride). Turn the gas to 3 psi. Be careful not to splash the extract out of the tube. <u>During concentration on the N-evap</u>, the internal wall of the concentrator tube and the N-evap sparging pipet must be rinsed down at least once or twice with ≈1 mL of methylene chloride. The solvent level in the concentrator tube must be positioned below the level of the water bath as much as possible to prevent water from condensing into the sample extract. As the extract volume is reduced, lower the N₂ sparging needle closer to the surface of the extract to expedite the concentration. Note any problems or extract losses, if they occur, in the extractions logbook.
- 7.37 Reduce each extract to slightly less than 1 mL and then, using a 5 ¾" pasteur pipet, transfer the final extract to a properly labeled 1.8 mL vial with PTFE-lined cap. Adjust the final volume of each extract to 1 mL using the 1 mL oil-filled reference vial for volume comparison. If the extract is highly colored or a precipitate forms during concentration, the final volume should be adjusted and noted in the extractions logbook with a comment.
- 7.38 Label the vial with lab sample number, extraction date, matrix and analysis. Store extract vials at a temperature of  $4 \pm 2$  °C until ready for analysis. Indicate in the extractions logbook the "tray location" of the individual extract vials.

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#### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

A method blank must be extracted for each and every item listed below:

- Each sample matrix (soil, water)
- Each day of extraction (24 hours midnight midnight)
- Each extraction method or level
- Every 20 samples extracted in a 24-hour period

A laboratory control sample (LCS) is required for <u>each</u> and <u>every</u> item listed below:

- Each sample matrix
- Each extraction method or level
- Every extraction batch of twenty or fewer samples
- 8.1 If a solvent blank or extraction blank is above the reporting level all associated samples with the "dirty" blank must be carefully evaluated versus the blank contamination level. Samples that contain DRO at a level of ten times or more than the blank level <u>may be useable</u>. If the sample level is less than ten times the blank level the source of the contamination is not as certain for the samples and they should be re-extracted after consultation with the client.
- 8.2 The analyst must make an initial demonstration of the ability to generate acceptable accuracy and precision with this method by successful analysis of the following:
  - 8.2.1 Replicate commercial diesel oil spikes in water: Analysis of at least 4 replicates at a concentration of 100  $\mu$ g/L (in water) with accuracy of the replicates falling between 60% to 140% of the known concentration. The precision of all replicates should be within 20%.
- 8.3 ME-DRO method LCS Requirements: every 20 water samples analyzed, the lab must analyze a set of duplicate diesel component spikes in reagent water. The Duplicate spikes must be run through the method in the same manner as samples. The accuracy of the two water spikes should fall between 60% to 140% of the known concentration with a relative % difference of 20% or less. Alternatively duplicate samples and spiked samples can be substituted for the laboratory spiked duplicates at a frequency of 10%. Care must be taken to ensure that the samples are homogeneous before analyzing duplicates and spikes.

Each extractions analyst must demonstrate proficiency in performing the extractions that prepare samples for analysis. Demonstration consists of preparation (extraction), by the analyst, of at least four aliquots which are then analyzed according to the analytical method in question. These QC samples must meet all quality control acceptance limits. Demonstration must be documented by use of a form which summarizes the results of the analysis of these aliquots, calculated percent recoveries, and standard deviation. Demonstration of proficiency

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must be done one time per analyst initially and then annually thereafter. Refer to SOPs QA-805 and QA-807, current revision.

If, upon analysis of the extracted samples, it is discovered that quality control acceptance criteria have not been met, all associated samples must be evaluated against all the QC. In some cases data may be reported, perhaps with narration, while in other cases, other corrective action may be taken. The corrective actions may include re-extraction of the samples associated with the quality control sample that did not meet acceptance criteria, or may include making new reagents and standards if the standardization is suspect. These decisions are based on holding time considerations, client and project specific Data Quality Objectives and on review of chromatograms. The supervisor, Operations Manager, and/or Quality Assurance Officer may be consulted to evaluate data. Some samples may not be able to be reanalyzed within hold time. In these cases "qualified" data with narration may be advisable after consultation with the client.

Much of the work performed at the lab is analyzed in accordance with specific QC requirements spelled out in a project specific Quality Assurance Project Plan (QAPP) or in a program specific Quality Systems Manual (QSM). The reporting limits, acceptance criteria and/or corrective actions may be different than those specified in this SOP. In these cases the appropriate information will be communicated to the Department Manager and/or senior chemists before initiation of the analyses so that specific product codes can be produced for the project. In addition, the work order notes for each project will describe the specific QAPP or QSM to be followed.

#### 9.0 METHOD PERFORMANCE

The method detection limit (MDL) is defined as the minimum concentration of a substance that can be measured and reported with 99% confidence that the value is above zero. The MDLs shall be determined and verified one time per type of instrument unless otherwise required by the method.

A Limit of Detection (LOD) is an estimate of the minimum amount of a substance that an analytical process can reliably detect. An LOD is analyte and matrix specific and may be laboratory-dependent. LODs must be determined for all parameters for which the laboratory is accredited under the DoD Environmental Laboratory Accreditation Program. LOD's must be verified for every preparation and analytical method combination and on every applicable instrument on a quarterly basis.

The Limit of Detection (LOQ) is the minimum levels, concentrations, or quantities of a target variable (e.g., target analyte) that can be reported with a specified degree of confidence. The LOQ shall be set at the lowest point in the calibration curve for all analyses utilizing an initial calibration. LOQ's must be verified quarterly for every preparation and analytical method combination and on every applicable instrument on a quarterly basis for all

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parameters included in the DoD Scope of Accreditation. The LOQ must be verified at least once annually if the analysis is not included in the DoD Scope of Accreditation.

MDLs are filed with the Organic Department Manager and then with the QAO. LOD and LOQ verifications are filed with the QAO

Refer to the current revision of Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, for procedures on determining the MDL.

Detection limit for waters: The laboratory must be able to achieve a detection limit of 50  $\mu$ g/L using a commercial diesel fuel oil mixture spiked into laboratory blank water and calculated against the DRO component standard.

Refer to the applicable analytical SOP for other method performance parameters and requirements.

#### 10.0 APPLICABLE DOCUMENTS/REFERENCES

ASTM "Std Mtds for Comparison of Waterborne Petroleum Oils by GC," 3328-78. Wisconsin DNR Modified DRO method, July 1993, Revision 6.

USEPA SW 846, 3rd edition, Methods 8000, 8100, 3500, 3510, 3520 and 3550.

The NELAC Institute, Laboratory Accreditation Standards, Volume 1, Management and Technical Requirements for Laboratories Performing Environmental Analysis, 10/06/2010.

Department of Defense Quality Systems Manual for Environmental Laboratories (DOD QSM), Current Version.

#### LIST OF TABLES AND FIGURES

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#### TABLE 1

#### QC REQUIREMENTS

Parameter/ Method	QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action
Sample Prep for waters and soils for DRO or TPH determination	Method blank	One per prep batch or 20 samples whichever is more frequent.	Refer to analytical method.	Refer to analytical method.
	LCS	One per prep batch	Refer to analytical method.	Refer to analytical method.
	Routinely  Matrix Spike/Matrix Spike Duplicate	One set for every 20 samples or 14 days whichever is more frequent, given sufficient sample volume.	Refer to analytical method.	Refer to analytical method.
	Per client Request Sample Duplicate/ Matrix Spike	One sample duplicate per twenty samples in conjunction with a matrix spike per 20 samples or 14 days whichever is more frequent.	Refer to analytical method.	Refer to analytical method.
	Demonstration of analyst proficiency; accuracy and precision	One time per analyst initially, and annually thereafter	Must pass all applicable QC for method	Repeat analysis until able to perform passing QC; document successful performance in personal training file
	MDL, LOD and LOQ studies and verifications			QA-806, Method Detection Limit rocedures on determining the

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# TITLE: PREPARATION OF AQUEOUS SAMPLES FOR ANALYSIS OF EXTRACTABLE PETROLEUM HYDROCARBONS or DIESEL RANGE ORGANICS (DRO)

#### TABLE 2

#### SUMMARY OF METHOD MODIFICATIONS

TOPIC	KATAHDIN SOP CA-520-07	METHOD
Apparatus/Materials Reagents	250 mL amber bottle used for extract collection     1 mL syringe     Short stem funnels	250 mL Erlenmeyer flasks     5 mL syringe     Drying columns
Sample preservation/ handling		
Procedures	<ol> <li>Extract collection in amber bottle or Erlenmeyer flask</li> <li>Add surrogate/spike to sample in CLLE</li> <li>Extract for 3 minutes on mechanical shaker</li> <li>Extract dried using Na<sub>2</sub>SO<sub>4</sub> in short stem funnels</li> <li>Rinse the extract flask three times with ~ 2 - 3 mLs of methylene chloride then rinse the sodium sulfate with ~ 15 mLs of methylene chloride to complete a quantitative transfer</li> <li>Water bath temp 75-85 deg C</li> <li>No apparatus height specification for concentration on water bath</li> <li>Sample removed from water bath when volume reaches ~6 mL</li> <li>N bath temp no higher than 39 deg C</li> </ol>	<ol> <li>Extract collection in Erlenmeyer flask</li> <li>Add surrogate/spike directly to sample bottle</li> <li>Extract by shaking vigorously for 1 - 2 minutes with periodic venting</li> <li>Extract dried using Na<sub>2</sub>SO<sub>4</sub> in drying columns</li> <li>Rinse the Erlenmeyer flask, which contained the solvent extract, with 20 - 30 mL of methylene chloride to complete the quantitative transfer</li> <li>Water bath temp 15-20 deg C above solvent boiling temp</li> <li>Partially immerse concentrator tube in water and lower apparatus to complete concentration in 10-20 min</li> <li>Sample removed from water bath when volume reaches 1 mL</li> <li>N bath temp 35 deg C</li> </ol>
QC - Spikes	o	o
QC - LCS	KAS does not perform.	Laboratory spiked duplicates prepared by spiking fuel oil into blank water in must be run at a minimum frequency of 5%. Alternatively duplicate samples and spiked samples can be substituted for the laboratory spiked duplicates at a frequency of 10%. Care must be taken to ensure that the samples are homogeneous before analyzing duplicates and spikes.
QC - Accuracy/Precision		
QC – MDL		

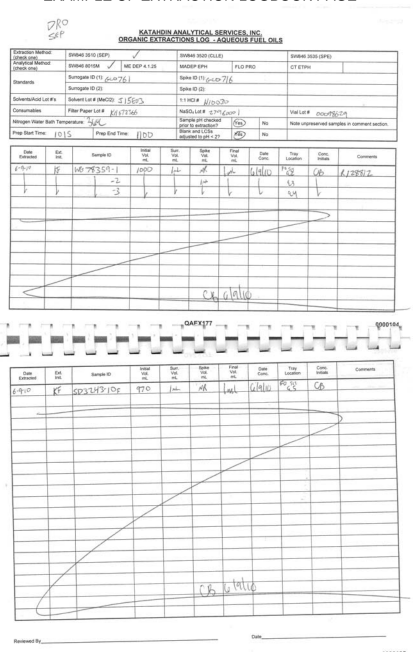
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# TITLE: PREPARATION OF AQUEOUS SAMPLES FOR ANALYSIS OF EXTRACTABLE PETROLEUM HYDROCARBONS or DIESEL RANGE ORGANICS (DRO)

FIGURE 1

EXAMPLE OF EXTRACTION LOGBOOK PAGE



# ADDENDUM SOP NO CHANGE FORM

# KATAHDIN ANALYTICAL SERVICES, INC. SOP "REVIEW WITH NO CHANGES" FORM

	5000 10 (1) (des
Name of Person Reviewing SOP: Jessica	Spear, VI-CO. C. C.
Review Date: 3-15-13	
SOP Number: CA - 520	
SOP Title: Preparation of aqueous of extractable petroleum N	s samples for analysis
THE ABOVE REFERENCED SOP HAS BEEN REVIE ANALYST OR SUPERVISOR. NO CHANGES ARE F	EWED BY A QUALIFIED AND TRAINED
Department Supervisor Signature:	Date:
Petro J	3-15-13
QAO Signature:	Date:
Le sa Dima ad	132113

# KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-333 Revision History Cover Page Page 1

		IATION OF PETI ENT OF ENVIRON			•	RO) BY FLORIDA FL-PRO
Prepared By	· -	Peter	Lemay		Date:	7/18/01
Approved By	<i>'</i> :		<i>(</i> )			
Group Super	visor:	- beter	Ly	·····	Date:	7/18/01
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QA Officer:	-	Detoah J.	nadeau		Date:	7.18.01
General Mar	nager:	Deina	uf. hu	Jehn .	Date:	7/18/01
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#### Revision History:

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
01	Added definitions and information for new data processing system. Added or challed word ingito clarify sections 6 and 7 and 7 able 2, Added wording to sections 8 and 9 per recent NELACH Navy audits. Minorchanges throughout. New Jigures 1 and 2.	- MRC	11,15,04	11,15.04
02	Changed Lims to Kims Sodium Sulfate is punied at vendor added wording to sect. 7.7.2 to elarify	LAO	03/06	03/06
03	Many changes made throughout including but not U miked to, waste management, CV brequency, Spike amounts, statistically derived Qr Limits and method modifications. Please refer to the DAM (SOP change form filed with the SOP in QA for a detailed list of ch	LAN	09/07	09/02
04	Removed Appartus and Reagend's that are not used.  Updated surrogate information	(An	09/08	09/08
	Sect. 8.4 and Table 1 - Changed Statistical Limits to method Limits. Added references. Updated Data Review Checklist	LAN	09/10	09/10

# KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-333 Revision History Cover Page (cont.)

Page 2

TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

Revision History (cont.):

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
06	8.6 — Changed CAR to nonconformance report. Section 9 - Added MDL/LOD/LOQ information. Updated Runlog and Review Checklist examples.	LAD	01/12	01/12
07	Sect. 1:7- Changed reporting from Quick- forms to Kins. Updated Fig. 2-Review Checklist. Minor edits.	LAD	05/13	05/13

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TITLE:	DETERMINATION OF PETROLEUM RANGE DEPARTMENT OF ENVIRONMENTAL PROT	
	acknowledge receipt of this standard operating preprovided. Return the bottom half of this sheet to the	
PETROL	owledge receipt of copy of document SOP DLEUM RANGE ORGANICS (PRO) BY DECTION METHOD # FL-PRO	CA-333-07, titled DETERMINATION OF DEPARTMENT OF ENVIRONMENTAL
Recipien	ent:	Date:
KATAHD	IDIN ANALYTICAL SERVICES, INC.	
	DARD OPERATING PROCEDURE	
PETROL	owledge receipt of copy of document SOP DLEUM RANGE ORGANICS (PRO) BY FLORIDA ECTION METHOD # FL-PRO	CA-333-07, titled DETERMINATION OF A DEPARTMENT OF ENVIRONMENTAL
Recipien	ent:	Date:

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### TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

#### 1.0 SCOPE AND APPLICATION

The purpose of this SOP is to describe the method used by Katahdin Analytical Services technical personnel to measure the concentration of petroleum range organics (PRO) in water and soil. These compounds correspond to a hydrocarbon range of  $C_8$ - $C_{40}$ .

This method is based on a solvent extraction, Gas Chromatography (GC) procedure. The method is designed to measure the petroleum concentration in environmental samples in the above stated C-Range (nominally diesel through motor oils). It cannot be used as an indication of gasoline contamination. Additional analyses may be performed including, but not limited to, analysis of additional reference materials. These additional efforts are not contained within this method.

#### 1.1 Definitions

PETROLEUM HYDROCARBONS: All chromatographic peaks, both resolved and unresolved, eluting between the peak start of n-octane (n- $C_8$ ) and the peak end after n-tetracontane (n- $C_{40}$ ). Quantitation is based on direct comparison of the area within this range to the total area of the Petroleum Hydrocarbon standard as determined from FID response using baseline-baseline integration.

PETROLEUM HYDROCARBON STANDARD: A 17-component mix of all even numbered normal alkanes from C8 to C40. This standard serves as a quantitation standard and a retention time window defining Petroleum Hydrocarbons.

SAMPLE MATRIX SPIKE: A selected sample from the analytical batch spiked with the Petroleum Hydrocarbon Standard and surrogate standards. The calculated spike recovery shall be used as a control.

LABORATORY CONTROL SAMPLE: Laboratory reagent grade water or standard soil spiked with the Petroleum Hydrocarbon standard and surrogate standards. The calculated spike recovery may be used as a laboratory control.

METHOD DETECTION LIMIT (MDL): Minimum concentration that an analyte can be measured and reported with a 99% confidence that the analyte concentration is greater than zero and is determined from analysis of a sample in a given matrix containing the analyte. The MDL is determined using EPA Appendix B to Part 136, CFR 40 Ch. 1(7-1-94) using the Student t Test.

KATAHDIN INFORMATION MANAGEMENT SYSTEM (KIMS): A complete multiuser system with the capabilities of integrating laboratory instrumentation, generating laboratory worksheets, providing complete Lab Order status and generating reports. LIMS utilizes these features through a database.

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### TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

PE NELSON TURBOCHROM: A data acquisition system that is used to collect chromatographic data. The system can also be used to archive raw data files.

TARGET: A software system that combines full processing, reporting and comprehensive review capabilities, regardless of chromatographic vendor and data type.

TARGET DB: An oracle database used to store and organize all Target data files.

#### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of analysts experienced in the analysis of PRO by Method FL-PRO. Analysts should be skilled in the interpretation of gas chromatograms and their use as a quantitative tool. Each analyst must demonstrate and document their ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, current revision, "Personnel Training & Documentation of Capability".

It is the responsibility of all Katahdin technical personnel involved in analysis of PRO by Method FL-PRO to read and understand this SOP, to adhere to the procedures outlined, and to properly document their data in the appropriate lab notebook. Any deviations from the test or irregularities with the samples should also be recorded in the lab notebook and reported to the Department Manager or designated qualified data reviewer responsible for this data.

It is the responsibility of the Department Manager to oversee that members of their group follow this SOP, to ensure that their work is properly documented and to initiate periodic review of the associated logbooks.

#### 1.3 Safety

Users of this procedure must be cognizant of inherent laboratory hazards, proper disposal procedures for contaminated materials and appropriate segregation of hazardous wastes. The toxicity or carcinogenicity of each reagent used in this method has not been precisely defined; however, each chemical should be treated as a potential health hazard. A reference file of material safety data sheets is available to all personnel involved in the chemical analysis. Everyone involved with the procedure must be familiar with the MSDSs (material safety data sheets) for all the materials used in this procedure.

Each qualified analyst or technician must be familiar with Katahdin Analytical Environmental Health and Safety Manual, including the Katahdin Hazardous Waste Management Plan and must follow appropriate procedures. These include the use of appropriate personal protective equipment (PPE) such as safety glasses, gloves and

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# TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

lab coats when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location of all safety equipment. Each analyst shall receive a safety orientation from their Department Manager, or designee, appropriate for the job functions they will perform.

#### 1.4 Pollution Prevention/Waste Disposal

Whenever possible, laboratory personnel should use pollution prevention techniques to address their waste generation. Refer to the current revision of the Katahdin Hazardous Waste Management Plan for further details on pollution prevention techniques.

Wastes generated during the preparation of samples must be disposed of in accordance with the Katahdin Hazardous Waste Management Plan and Safety Manual and SOP SD-903, "Sample Disposal," current revision. Expired standards are lab packed, placed in the Katahdin hazardous waste storage area, and disposed of in accordance with this SOP.

Any methylene chloride solvent waste generated during the preparation of standards etc. should be disposed of in the "D" waste stream satellite accumulation area nearest the point of generation. Acetone and hexane are considered flammable waste, and should be disposed of in the "O" waste stream satellite accumulation area nearest the point of generation. FLPRO sample vials are considered "P" waste and should be disposed of in the corresponding satellite waste accumulation area nearest the point of generation. Please refer to the current revision of SOP CA-107 for the location of satellite waste accumulation areas.

#### 2.0 SUMMARY OF METHOD

- One liter of water or a specified quantity of soil (extraction method dependent) is spiked with two surrogates and extracted with Methylene chloride. The water is removed from the extract, concentrated to a volume of 2.0 mL, and treated with silica gel to remove potential organic interferences. An aliquot is injected onto a capillary column gas chromatograph (GC) equipped with a flame ionization detector (FID). Quanitation is based on the detector response compared to a series of normal alkane standards. This method is suitable for the analysis of waters, soils or wastes.
- 2.2 This method is based in part on USEPA Methods 8000 and 8100, SW-846, "Test Methods for Evaluating Solid Waste", 3<sup>rd</sup> Edition, Method OA-2, work by the EPA UST Work Group "Measurement of Petroleum Hydrocarbons: Report on activities to Develop a Manual", 1990, Method AK103.0, Revision 2, PUBL-SW-141, July 1993

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# TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

and the Florida Department of Environmental Protection Technical Advisory Committee for 62-770, F. A. C, Petroleum Contamination Site Cleanup Criteria.

#### 3.0 INTERFERENCES

- 3.1 Other organic compounds including chlorinated hydrocarbons, phenols, and phthalate esters are measurable. As defined in the method, the PRO results include these compounds. Spills of known specific constituents should be analyzed and quantified by a method specific for those compounds.
- 3.2 Method interferences are reduced by washing all glassware with hot soapy water and then rinsing it sequentially with tap water, methanol or acetone, and Methylene chloride. Method blanks must be analyzed with each batch to demonstrate that the samples are free from method interferences.
- 3.3 High purity reagents (pesticide grade or better) must be used to minimize interferences.
- 3.4 Contamination by carryover can occur whenever high-level and low-level samples are sequentially analyzed. Whenever an unusually concentrated sample is encountered, it should be followed by analysis of a solvent blank to check for cross-contamination.
- 3.5 Animal and vegetable oil and grease and biogenic terpenes are also measurable if the sample is not cleaned up before analysis. In order to eliminate false positives from these sources, the silica cleanup is a mandatory part of the procedure.

#### 4.0 APPARATUS AND MATERIALS

- 4.1 Gas Chromatograph: Analytical system complete with gas chromatograph and all required accessories, including a detector, column supplies, recorder, gases, and syringes. A capillary split/splitless injector operating in the splitless mode is recommended. A data system capable of determining peak areas by integrating from baseline to baseline is required.
- 4.2 Column 1: 30 m x 0.53 mm ID ZB-5, 1.5 micron film thickness (or equivalent). Column 2: 30 m x 0.53 mm ID ZB-1, 1.5 micron film thickness (or equivalent). The column must be capable of resolving typical diesel components, and the solvent front from  $C_8$ . Other columns may be used if all column performance criteria are met.
- 4.3 Detector: Flame ionization detector (FID).

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### TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

- 4.4 Microsyringes: 1 ul, 5 ul, 10 ul, 25 ul, and 100 ul.
- 4.5 Disposable pipettes: Pasteur.
- 4.6 2 ml (and larger) vials with Teflon lined caps for storage of extracts.

#### 5.0 REAGENTS

- 5.1 Solvents: Methylene chloride: Pesticide grade or equivalent. Store away from other solvents.
- 5.2 Stock Standards: Aliphatic Hydrocarbon standard mix from a vendor like UltraScientific at a concentration of 500 ug/mL in hexane (each of the 17 components from C<sub>8</sub> to C<sub>40</sub>). A surrogate solution containing n-Triacontane-d<sub>62</sub> at a concentration of 5000 ug/mL and another surrogate solution containing o-Terphenyl at a concentration of 2000 ug/mL from a vendor like Restek.
- 5.3 Calibration Standards: The standards are prepared at the following five different concentrations: 200 ug/ml, 100 ug/ml, 50 ug/ml, 20 ug/ml, and 5 ug/ml (per each component). This is equivalent to 85, 340, 850, 1700, and 3400 ug/ml total alkanes in the standards. The concentration of OTP and triacontane-d<sub>62</sub> must remain at a constant 50 and 300 ug/ml level in all concentration levels.
  - 5.3.1 Transfer the stock standard solution into a Teflon-sealed screw-cap/crimp cap bottle. Store, with minimal headspace, at 6°C or less and protect from light.
  - 5.3.2 Working standards must be replaced after 6 months, or sooner if comparison with check standards indicates a problem.

#### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

- 6.1 Whenever possible, samples should be grab samples which are collected directly into the sample container. Sample collection equipment such as bailer or intermediate containers should be avoided (exceptions: collection from monitoring wells or grab samples in surface water at depth). Unless required by permit, automatic samplers may not be used. Pumps such as bladder pumps or peristaltic pumps shall not be used.
- 6.2 All sampling equipment which contacts the sample shall be constructed of teflon®, stainless steel or glass. Under no circumstances can flexible PVC tubing, such as tygon®, be used in the purging or sample collection process.

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# TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

6.3 Water samples shall be collected in a one liter glass container; soils in a glass jar. All containers shall be sealed with a screw cap with teflon® liner. Water samples shall be acidified to a pH of less than 2 with hydrochloric or sulfuric acid (reagent grade or better).

6.4 The samples shall be stored at 4°C (±2°C) from the time of collection until extraction. Extraction shall be performed on waters within seven days of sample collection and on soils within 14 days of sample collection. All analyses must take place within 40 days of extraction.

#### 7.0 PROCEDURES

7.1 Waters are extracted using a separatory funnel or continuous liquid liquid extraction technique. Soils are extracted using a sonication technique. Alternatively, soils may be extracted by a Soxhlet extraction technique. Refer to Katahdin SOP CA-520, current revision, for sample preparation procedures. After the extracts are concentrated, an appropriate volume (usually 1ul) is injected directly into the GC. (Recommend using splitless injection techniques).

NOTE: NaCl may be added to water samples to improve extraction efficiency.

If the sample concentration exceeds the calibration range for PRO an appropriate dilution should be used. An appropriate dilution is one that keeps the response of major constituents (previously saturated peaks) in the linear range of the detector. If an initial dilution does not accomplish this then an intermediate dilution should be performed.

#### 7.2 Gas Chromatography:

- 7.2.1 Conditions (For both column 1 and 2): Set column temperature to 60°C for 2 minutes, then 10°C/min. to 300°C and hold for 24 min. Set FID Detector to 310°C and injector to 300°C. Conditions may be altered to improve resolution or recovery of petroleum range organics.
- 7.2.2 Performance Criteria: GC run conditions and columns must be chosen to meet the following criteria:
  - 7.2.2.1 Resolution of C<sub>8</sub> from the solvent front.
  - 7.2.2.2 The column must be capable of separating typical petroleum hydrocarbon components from the surrogates.

#### 7.3 Retention Time Window

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# TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

- 7.3.1 Before establishing windows, be certain that the GC system is within optimum operating conditions. Make three injections of the method standard throughout the course of a 72 hour period. Serial injections over less than a 72 hour period result in retention time windows that are too tight.
- 7.3.2 Calculate the standard deviation of the absolute retention times for the two surrogates,  $C_8$ , and  $C_{40}$ .
  - 7.3.2.1 The retention time window for individual peaks is defined as a plus or minus three times the standard deviation of the absolute retention time for each component.
  - 7.3.2.2 In those cases where the standard deviation for a particular analyte is zero, the laboratory should use  $\pm 0.05$  min as a retention time window.
- 7.3.3 The laboratory must calculate retention time windows for these standards on each GC column and whenever a new GC column is installed. The data are retained by the laboratory.

#### 7.4 PRO Calibration

7.4.1 Initial Calibration – Calibration shall be by external calibration using a minimum of 5 concentration levels for the initial calibration. Quantitation shall be by linear regression.

In all cases, response of the standards must be determined by continuous integration of all responses (excluding surrogates) from a forced baseline beginning at a point prior to the elution of  $C_8$  to a point past  $C_{40}$ . All responses must be determined as responses to baseline and not valley to valley. A method is calibrated for all five levels using the area of each of the 17 individual alkanes and the area of the two surrogates and a total area of the Petroleum Hydrocarbon Standard (PRO) (which is the total area of the seventeen alkanes for each level).

- 7.4.1.1 Linear Regression The linear regression shall be calculated using the total PRO area versus the PRO concentration. The correlation coefficient shall be equal to or greater than 0.995.
- 7.4.1.2 The accuracy of the initial calibration shall be verified by injecting a midpoint concentration of a standard mix that has been obtained from a different source. The calculated value shall be within  $\pm$  20% of the expected value.

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# TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

7.4.2 Continuing Calibration – The calibration curve must be verified on each working day by the injection of a continuing calibration standard (CV) at a midpoint concentration. This standard must be evaluated prior to the analysis of samples.

In addition, a continuing calibration must be run every 10 samples and at the end of the sequence. The concentration of these should vary, with at least one at a level of 1-2 times the calculated PQL as a verification of sensitivity. To accomplish this, continuing calibrations at 50 ug/ml and 20 ug/ml (each component) should be ran.

- 7.4.2.1 If the concentration of this standard varies from the predicted concentration by more than  $\pm$  25%, a new initial calibration curve must be prepared and verified before samples are analyzed.
- 7.4.2.2 Retention Time Window Establish daily retention time windows for each analyte of interest using the absolute retention time for each analyte as the midpoint of the window for that day **if** after analyzing the midpoint it is determined that one or more analytes falls outside of the previously established absolute retention time window. The daily retention time window equals the midpoint ± three times the standard deviation determined in Section 7.3.
- 7.5 Gas Chromatography Analysis
  - 7.5.1 A 1 ul injection volume is analyzed by GC/FID.
  - 7.5.2 If an initial calibration has already been performed, verify the calibration by analysis and evaluation of a mid-point CV on each working day.
    - In addition, a CV must be run every 10 samples and at the end of the sequence.
  - 7.5.3 Evaluate the CV per 7.4.2.1 and 7.4.2.2. If either performance criteria fails, the instrument must be recalibrated and all samples which were injected after the failed standard must be reanalyzed.
  - 7.5.4 A Methylene chloride blank will be run in every sequence to determine the area generated on normal baseline bleed under the conditions prevailing in the 24 hour period if requested by the client. This area is determined by continuous integration of all responses under the same conditions (i.e. forced baseline and predetermined time interval) as the samples. This blank is calculated as the solvent blank and the value should be less than the PQL.

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# TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

Methylene chloride blanks should also be run after samples suspected of being highly concentrated to prevent carryover. If the blank analysis shows contamination, additional blanks should be analyzed until the system is shown to be free from contaminants.

- 7.5.5 If the sample concentration exceeds the linear range of the method in the final extract, the extract must be diluted and reanalyzed.
- 7.5.6 Baseline correction is allowed to correct for rises due to temperature programming. Range integration is corrected by the automatic subtraction of the baseline established by activation of a programmed run without the injection of any material. Instrument baseline must be established for every batch of samples.

#### 7.6 Calculations

- 7.6.1 The integrated area for all peaks eluting from n-octane through n-tetracontane shall be determined using a baseline drawn from the baseline point to n-octane to a point past n-tetracontane where the baseline returns to normal. All area including the "hump-a-gram" and surrogate standards shall be included. Do not integrate valley to valley for individual peaks except for the two surrogates. The concentration of the PRO is calculated by using the calibrated curve that is prepared in Target. Target displays a concentration when the file is processed through the appropriate calibrated method.
- 7.6.2 The concentrations from the reports are then incorporated with the extraction data to arrive at a final concentration.

7.6.2.1 Water: Conc (ug/L) = (Amt) (DF) ((Vt/Vo) 1000)

7.6.2.2 Soil/Sediment: Conc (mg/kg) = (Amt) (DF) ((Vt/Vo) (100/(100-M)))

where, Amt = adjusted concentration calculated by Target in ug/ml

Vt = Volume of total extract

Vo = Volume or weight of sample extracted

M = % Moisture
DF = Dilution Factor

#### 7.7 Data Review

#### 7.7.1 Initial Data Review

The initial data review is accomplished by the analyst who ran the samples. This review is of sufficient quality and detail to provide a list of samples that

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# TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

need to be reanalyzed or diluted and reanalyzed. The initial data review is performed in Target Review. This data review examines criteria that directly impact whether or not the sample needs to be reanalyzed and/or extracted. These criteria include:

- ◆ QC criteria for method blank, LCS, MS/MSD, and calibration refer to section 8.0.
- Surrogate recovery
- Chromatography: manual integration.
- Target compound detection: quantitation, false positives.

The requirement of the GC laboratory is that this initial data review be completed no later than the end of the next work day. After the analyst has completed his or her initial data review, the information is then ready to be processed for reporting. Refer to section 7.8.

#### 7.7.2 Surrogate recovery

The recoveries for o-Terphenyl are compared to the method acceptance limits. The recoveries for n-Triacontane-d<sub>62</sub> are compared to nominal acceptance limits of 70-130% until laboratory acceptance limits can be established.

The sample is evaluated for recoveries of the surrogate OTP and n-Triacontane- $d_{62}$ . If the recovery is low and there is no apparent matrix effect, the sample should be reanalyzed. If the reanalysis is still low, re-extract. If the recovery is low and there may be a matrix effect, reanalyze to confirm a matrix effect and narrate. If the surrogate is high and the sample results are less than the PQL, or there is likely a matrix effect, narrate.

#### 7.7.3 Chromatography

The chromatography should be examined for the presence of any non-target peaks, which can be used as an indication of whether or not matrix interference might be influencing surrogate recoveries.

Manual integrations are to be performed when chromatographic conditions preclude the computer algorithm from correctly integrating the peak of concern. In no instance shall a manual integration be performed solely to bring a peak within criteria.

In Target Review, each peak of concern is examined by the primary analyst to ensure that the peak was integrated properly by the computer algorithm. Should a manual integration be necessary (for instance, if the sample contains a concentration of PRO which was integrated "valley to valley"

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# TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

instead of a "baseline to baseline"), manual integration is performed in Target Review. A "m" qualifier will automatically be printed on the quantitation report summary indicating that a manual integration was performed. For specific procedures on how to manually integrate, refer to Katahdin SOP QA-812, Manual Integration, current revision.

#### 7.8 Reporting

After the chromatograms have been reviewed and any target analytes have been quantitated using Target, the necessary files are brought into KIMS. Depending on the QC level requested by the client, a Report of Analysis (ROA) and additional reports, such as LCS forms and chronology forms, are generated. The package is assembled to include the necessary forms and raw data. The data package is reviewed by the primary analyst and then forwarded to the secondary reviewer. The secondary reviewer validates the data and checks the package for any errors. When completed, the package is sent to the Department Manager for final review. A completed review checklist (Figure 2) is provided with each package. The final data package from the Organics department is then processed by the Data Management department.

#### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

See below or refer to Table 1 for a summary of QC requirements, acceptance criteria, and corrective actions. These criteria are intended to be guidelines for analysts. The table does not cover all possible situations. If any of the QC requirements are outside the recovery ranges listed in Table 1, all associated samples must be evaluated against all the QC. In some cases data may be reported, but may be reanalyzed in other cases. Making new reagents and standards may be necessary if the standardization is suspect. The corrective actions listed in Table 1 may rely on analyst experience to make sound scientific judgments. These decisions are based on holding time considerations, client and project specific Data Quality Objectives and on review of chromatograms. The Department Manager, Operations Manager, and/or Quality Assurance Officer may be consulted to evaluate data. It may not be possible to reanalyze samples within hold time. In these cases "qualified" data with narration may be advisable after consultation with the client.

In some cases the standard QC requirements listed in this section and in Table 1 may not be sufficient to meet the Data Quality Objectives of the specific project. Much of the work performed at the lab is analyzed in accordance with specific QC requirements spelled out in a project specific Quality Assurance Project Plan (QAPP) or in a program specific Quality Systems Manual (QSM). The reporting limits, acceptance criteria and/or corrective actions may be different than those specified in this SOP. In these cases the appropriate information will be communicated to the Department Manager and/or senior chemists before initiation of

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the analyses so that specific product codes can be produced for the project. In addition, the work order notes for each project will describe the specific QAPP or QSM to be followed.

- 8.1 For each analytical batch (up to 20 samples), a method blank, laboratory control sample (LCS), matrix spike and matrix spike duplicate or sample duplicate are analyzed. They are carried through all stages of the sample preparation and analysis steps.
- 8.2 The laboratory shall generate control limits based on +/-3 standard deviations from the average recovery for all spikes and surrogates, and + 3 standard deviations from the average precision value for all duplicates. The limits that area generated must be within the criteria specified in Table 3 below.
- 8.3 Spike concentrations: The LCS and the MS/MSD are spiked with the seventeen component PRO mix at the same concentration. The spike concentrations are:

	WATER ug/L	SOILS mg/Kg
PRO	850	28.5

The surrogate spike concentrations in the final extract are:

	WATER	SOILS
	ug/ml	mg/kg
o-Terphenyl	50	1.65
n-Triacontane-d <sub>62</sub>	300	10

8.4 LCS and MS/MSD acceptance criteria and Corrective Action: All QC samples are calculated for percent recovery of the spiked analyte(s). The recoveries are compared to the method acceptance limits. Refer to Table 3 for these limits.

If any spike compound in the laboratory control sample falls outside of the established recovery acceptance limit window, the QC sample is considered to be out of control and any sample that is associated should be reextracted. However, if the recovery is high and the associated samples do not contain the specific compound(s), the data can possibly be accepted with narration.

If a spike compound is outside of the acceptance limits in the matrix spike sample but is acceptable in the LCS, the data is considered acceptable. The cause of the failure is possibly attributable to matrix interference. However, if the compound fails in both the LCS and the MS/MSD, the result for that analyte is suspect and may not be reported for regulatory compliance purposes.

8.5 Surrogate acceptance criteria and Corrective Action: Surrogate recoveries are calculated on all samples, blanks and spikes. The recoveries for o-Terphenyl are compared to the method acceptance limits. The recoveries for n-Triacontane-d<sub>62</sub> are

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compared to nominal acceptance limits of 70-130% until laboratory acceptance limits can be established.

When a sample has a surrogate that falls outside of the method acceptance limit window, the problem should be investigated. If the recovery looks like it is affected by the sample matrix, the sample may be reinjected to confirm matrix interference. When a sample has no detectable surrogate recovery, the sample should be reextracted

8.6 Non-conformance Report: Whenever data is not acceptable because of a failing LCS or surrogate recovery, a non-conformance report (NCR) must be initiated as soon as possible.

#### 9.0 METHOD PERFORMANCE

- 9.1 The MDL of this method is estimated to be at least 4 mg/kg for soil and 0.1 mg/L for water. Each laboratory shall establish a laboratory specific MDL for all matrices prior to analyzing any samples.
- 9.2 The method detection limit (MDL) is defined as the minimum concentration of a substance that can be measured and reported with 99% confidence that the value is above zero. The MDLs shall be determined and verified one time per type of instrument unless otherwise required by the method.
- 9.3 Limits of Detection (LOD) is an estimate of the minimum amount of a substance that an analytical process can reliably detect. An LOD is analyte and matrix specific and may be laboratory-dependent. LODs must be determined for all parameters for which the laboratory is accredited under the DoD Environmental Laboratory Accreditation Program. LOD's must be verified for every preparation and analytical method combination and on every applicable instrument on a quarterly basis.
- 9.4 The Limit of Quantitation (LOQ) is the minimum levels, concentrations, or quantities of a target variable (e.g., target analyte) that can be reported with a specified degree of confidence. The LOQ shall be set at the lowest point in the calibration curve for all analyses utilizing an initial calibration. LOQ's must be verified quarterly for every preparation and analytical method combination and on every applicable instrument on a quarterly basis for all parameters included in the DoD Scope of Accreditation. The LOQ must be verified at least once annually if the analysis is not included in the DoD Scope of Accreditation.
- 9.5 MDLs are filed with the Organic Department Manager and then with the QAO. LOD and LOQ verifications are filed with the QAO

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# TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

- 9.6 Refer to the current revision of Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, for procedures on determining the MDL.
- 9.7 Refer to the current revision of the Florida Department of Environmental Protection Method for Determination of Petroleum Range Organics (Method # FL-PRO) for other method performance parameters and requirements.

#### 10.0 APPLICABLE DOCUMENTS/REFERENCES

Florida Department of Environmental Protection, Method for Determination of Petroleum Range Organics, Method # FL-PRO, Revision 1, November, 1995.

ASTM "Standards Methods for Comparison of Waterborne Petroleum Oils by Gas Chromatography," 3328-78.

Wisconsin DNR Modified DRO method, July 1993, Revision 6.

Department of Defense Quality Systems Manual for Environmental Laboratories (DOD QSM), Current version.

The National Environmental Laboratory Accreditation Conference (NELAC) Standards, June 2003.

The NELAC Institute, Laboratory Accreditation Standards, Volume 1, Management and Technical Requirements for Laboratories Performing Environmental Analysis, 10/06/2010.

Katahdin SOP CA-101, Equipment Maintenance and Troubleshooting, current revision.

Katahdin SOP QA-803, Laboratory QA: Self Inspection System, current revision.

Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, current revision.

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#### LIST OF TABLES AND FIGURES

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Table 2	Summary of Method Modifications
Table 3	Method Acceptance Criteria
Table 4	PQLs
Figure 1	Instrument Runlog Page
Figure 2	Data Review Checklist

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# TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

#### TABLE 1

#### QC REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action
Method blank	One per prep batch	No analyte detected >PQL	(1) Investigate source of contamination (2) Evaluate the samples and associated QC: i.e. If the blank results are above the PQL, report sample results which are < PQL or > 10X the blank concentration. Otherwise, reprep a blank and the remaining samples.
LCS	One per prep batch	Method acceptance limits	(1) Evaluate the samples and associated QC: i.e. If an MS/MSD was performed and acceptable, narrate. If an LCS/LCSD was performed and only one of the set was unacceptable, narrate. If the surrogate recoveries in the LCS are low but are acceptable in the blank and samples, narrate. If the LCS recovery is high but the sample results are < PQL, narrate. Otherwise, reprep a blank and the remaining samples.
Initial Calibration	Initial cal prior to sample analysis	Correlation coefficient => 0.995	(1) Perform instrument maintenance as needed. (2) Reanalyze and or reprep calibration standards.
CV( At or near the midpoint of the ICAL)	On each working day prior to sample analysis if an ICAL was previously analyzed	± 25 %D	(1) Evaluate the samples: If the %D>+25% and sample results are <pql, %d="" if="" narrate.="">±25% and is likely a result of matrix interference, narrate. All samples must be reanalyzed that fall within the standard that exceeded criteria and the last standard that was acceptable.</pql,>
End of sequence CV	At the end of each 12-hour work shift or after running 10 samples, whichever is sooner	± 25 %D	<ul><li>(1) Evaluate sample data if criteria exceeded due to matrix; narrate, and perform maintenance for new samples.</li><li>(2) If criteria are exceeded and this is not due to matrix, Reanalyze.</li></ul>
Matrix Spike/Matrix Spike Duplicate	One for every set of 20 samples provided samples aliquots are not depleted	Laboratory established acceptance limits RPD< 20 % for waters and < 25 % for solids	(1) Evaluate the samples and associated QC: i.e. If the LCS results are acceptable, narrate. (2) If both the LCS and MS/MSD are unacceptable, reprep the samples and QC.
Sample Duplicate ( If required in lieu of MSD)	One sample duplicate per twenty samples	RPD ≤20 for waters, RPD ≤25 for solids	(1) Evaluate data for matrix interference homogeneity of sample.
Demonstration of analyst proficiency; accuracy and precision	One time per analyst initially and annually thereafter	Must pass all applicable QC for method	Repeat analysis until able to perform passing QC; document successful performance in personal training file

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### TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

TABLE 1 (cont.)

#### QC REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action		
Demonstration of analyst proficiency; accuracy and precision	One time per analyst initially and annually thereafter	Must pass all applicable QC for method	Repeat analysis until able to perform passing QC; document successful performance in personal training file		
MDL study/ LOD, LOQ Verifications		efer to KAS SOP QA-806, "Method Detection Limit, Instrument Detection Limit and Reporting mit Studies and Verifications", current revision.			

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## TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

#### TABLE 2

#### SUMMARY OF METHOD MODIFICATIONS

TOPIC	KATAHDIN SOP CA-333-07	FL-PRO, current revision
Apparatus/Materials		
Reagents	5.3 Sodium sulfate purified by vendor     5.5 PRO free sand ( Muffled )	7.3 Sodium sulfate purified by heating at 400 deg C for 4 hours or extracting x3 with methylene chloride and drying at 105 deg C. 3.4 Ottawa sand
Sample preservation/ handling		
QC – Method Blank	Table 1 No analyte detected >PQL	10.4TRPH value of the blank shall be at or below the established method detection limit.
QC - Surrogates	Use n-Triacontane-d <sub>62</sub> . Use nominal limits of 70-130 until laboratory limits can be established.	7.4.1 Recommend C <sub>39.</sub> Use method acceptance limits.
QC - Spikes	8.2PRO concentration of 850 ug/L for waters and 28.5 mg/kg for soils.	7.4.4 Total PHS concentration in the spiked sample of 5 mg/L in water or 300 mg/kg in soilsThe concentration of the spike in the sample should be approximately 3-5 times the level expected in the samplelevel of the spike should be adjusted
QC - Accuracy/Precision		
QC - LCS		
QC - MDL		
Procedure	7.4.2.2 Retention Time Window – Establish daily retention time windows for each analyte of interest using the absolute retention time for each analyte as the midpoint of the window for that day if after analyzing the midpoint it is determined that one or more analytes falls outside of the previously established absolute retention time window. The daily retention time window equals the midpoint ± three times the standard deviation determined in Section 7.3.	9.3.2.2 Retention Time Window – The retention time window for the surrogates and C8 and C40 shall be within the established range If they are out of acceptance range, a new initial calibration must be prepared and verified before samples are analyzed.
	7.5.4 The Methylene chloride blank is analyzed if requested by the client and will be less than the PQL.	9.5.4 The methylene chloride blank must be analyzed with each sequence and the PRO concentration shall be less than the MDL of the method.
	7.5.4 Baseline correction is allowed to correct for rises due to temperature programming. Range integration is corrected by the automatic subtraction of the baseline established by activation of a programmed run without the injection of any material. Instrument baseline must be established for every batch of samples.	9.5.4 Do not baseline subtract  7.4.3 Suggested calibration levels are 5, 50, 150,
	5.7 The standards are prepared at the following five different concentrations: 200 ug/ml, 100 ug/ml, 50 ug/ml, 20 ug/ml, and 5 ug/ml (per each component).	250, 350 and 500 ug/mL of each individual component.

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### TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

### TABLE 3

#### % Recovery Precision (%RSD) Water Soil Water Soil Soxhlet **Sonication** Sonicati Soxhlet on Matrix Spike Samples 41-101 41-224 62-204 0-20 0-25 0-25 Laboratory Control Spike Samples 63-135 55-118 63-153 0-20 0-25 0-25 Surrogates: OTP 82-142 57-115 62-109 n-triacontane-D<sub>62</sub> 70-130 70-130 70-130

METHOD ACCEPTANCE CRITERIA

TABLE 4

#### **PQLS**

Analyte	Water ug/L	Soil mg/Kg
PRO	500	20

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#### FIGURE 1

#### EXAMPLE OF RUNLOG PAGE

		d	1	Metho	od (circle): EPH(M	ADEP) (F	EL PRO JTNRCC 1005
Reviewe	d by/ [	Date:		DRO/	TPH - 8015Mod.	/ MDEP 4	.1.25 / 8100Mod.
Date	Init.	Result File	Sample ID	Y/N	Method	Column	Comments
5-5-11	AC	CEE2012	TB	N	AROBO31A	366	
1		1 13		1		1	
		14					
		15	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	1			
		16	SE 2262-1	Y			
		17	M	N			
		18		11			
_		19					
-		20	-	1			
-	-	21	SE2316-1	y			
-	+	22	TB	N			
-		23	l l	Y			
	1	24	AR050	y	, ,		Hos8
11	0/	1	rever Lin	1	Septa		
5-5-11	AC	25	TB -0	N.	FLPBOZIA		
1	++	26	TB FLP50	1			
	+	28		V			H2009
+	+	29	1 200	1			Hze11
++	+	30	100	V			H2012
++	+	31	5	Y			H 700'
	+	32	IND	Y			H2013
	+	33	TB	V			H2014
1	++		WG91154-1	N		<b>.</b>	
6-11	1	35	1 -2	N			Α
1	1	36	1 -3	N	-		spike1
	1		AQ LOD	V			1
	11	38	LOG	4			
	1		5E2433-2	N			
1	1	1 40	1 -3	N			554

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### TITLE: DETERMINATION OF PETROLEUM RANGE ORGANICS (PRO) BY FLORIDA DEPARTMENT OF ENVIRONMENTAL PROTECTION METHOD # FL-PRO

#### FIGURE 2

#### EXAMPLE OF DATA REVIEW CHECKLIST

#### PRIMARY REVIEW CHECKLIST

ent:	Primary	Secon	dary
thod:	Date:	Date:	
G No: Level:	Initials:	Initials:	
S No:		Approved :	□ Y
DODOGW (44) D. DOD W	/ LAD LIMITS -	NUADD T	4 D 🗆
DODQSM (4.1) $\square$ DOD W. (REPORT ND's to		QUAPP □ L. LOD □)	AB □
List all curves that are scanned (h	ard copy not included ).		
Narrate which QC limits were us	ed for ( Surr., LCS's MS/MS	D's.)	
All needed forms are present.			
Correct Work Order Number or S	DG name (all forms).		
Correct project name and spelling	(all forms). (Truncated $\square$ )	,	
Correct file numbers (all forms).			
Analysis Date Correct.			
Extraction Method & Analysis Me	ethod Correct.		
Product list compared to ROAs (c	ompounds & PQLs).		
Chromatogram reviewed for unlab	peled peaks (check product li	st).	
Flagging of all ROAs correct (Fl	orida 🗆 ) ( Florida 🗆 ).		
All tunes included (level IV).			
All log book pages included (Soil	weights, TCLP & SPLP).		
Verify DOD QSM criteria and/or	Project specific requirement	s.	
Narrate any method deviations.	(Blanks, LCS's etc. )		
Sign & Date Manual integration (	Narrate as needed ).		
	ATED). YES Ple		ow:

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### KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-316 Revision History Cover Page Page 1

TITLE: METHOD FOR DETERMINING VOLATILE PETROLEUM HYDROCARBONS or GASOLINE RANGE ORGANICS (GRO) BY METHOD 8015 (WITH OR WITHOUT MODIFICATIONS)

Prepared By:	Peter Lemay	Date:	6/97
Approved By:			
Group Supervisor:	betwo Ly	Date:	5/24/01
Operations Manager:	Schutz	Date:	5/23/01
QA Officer:	actoral J. Nadeau	Date:	5.24.01
General Manager:	Dernau F. Kulgalu	Date:	924/01
			′ /

#### Revision History:

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
02 8015B	Revised Sections 7.3.2, 7.4.4, 7.4.1, and added 5.9 to reflect the option of using method sois with no modifications—i.e. for South Carolina	9n	52401	5.24.01
	Minor editorial changes to sec-			
03	tions 7.7.3 and 9.0.	2n	5.22.02	5:22:07
80153		,		
04	Added definitions and information			
8015B	for new data processing system. Added or changed wording to clarify sections 8+9. Minor changes throughout.	HRC	11.09.04	11.09.04
<b>೧</b> ೮	removed references to TFT	1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -		
	added atternate levels for CV's removed Sect. 8.13 Cgasoline composite sample)	LAN	04/00	04/06
801503				
06	Removed all remaining references to TFT and extraction surrogates. Changed wording in Sect. 7.4. to reflect corrent code bration procedure. Added wording to Sect. 7.9.4 clarifying BFB. J. GRO quantition. Added wording to sect. 8.1.2 darking sample regulas. Added ICV information to Sect. 1.5.7 and Tables	100	07/07	07/07

SOP Number: CA-316 Revision History Cover Page Cont. Page 2

TITLE: METHOD FOR DETERMINING VOLATILE PETROLEUM HYDROCARBONS or GASOLINE RANGE ORGANICS (GRO) BY METHOD 8015 (WITH OR WITHOUT MODIFICATIONS)

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
07	Added use of Casoline Standard for fingerprinting. Minor formatting Changes throughout. Edited Section numbers. Clarified Correlation coefficient Criteria. Added figure for GC Soil prep logbook.	UAD	03/08	03/08
08	Added interference regarding FID. Updated method references. Updated Method Medification Table 2.	LAN	0 2/09	०२०९
09	update RT window criteria, method bleak criteria and "Q" plagging criteria for compliance with DoD QSM version 4.1. Added references.	LAD	08109	08/09
10	Added Chemstation Lifiation. Ghanged QC Limits to Laboratory Statistically derived acceptance Limits. Added LOQ, LOD, & MDL informa Updated and added references Updated Figure		03/12	03/12

KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-316-10 Date Issued: 03/12

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TITLE:		MINING VOLATILE PETROLEUM HYDROCARBONS or GANICS (GRO) BY METHOD 8015 (WITH OR WITHOUT
		candard operating procedure by signing and dating both of the half of this sheet to the QA Department.
DETERM	IINING VOLATILE PETROLI	of document SOP CA-316-10, titled METHOD FOR EUM HYDROCARBONS or GASOLINE RANGE ORGANICS WITHOUT MODIFICATIONS).
Recipien	t:	Date:
	IN ANALYTICAL SERVICES RD OPERATING PROCEDU	,
DETERM	IINING VOLATILE PETROL	of document SOP CA-316-10, titled METHOD FOR EUM HYDROCARBONS or GASOLINE RANGE ORGANICS WITHOUT MODIFICATIONS).
Recipien	<b>.</b> .	Date:

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TITLE: METHOD FOR DETERMINING VOLATILE PETROLEUM HYDROCARBONS or GASOLINE RANGE ORGANICS (GRO) BY METHOD 8015 (WITH OR WITHOUT

**MODIFICATIONS**)

#### 1.0 SCOPE AND APPLICATION

The purpose of this SOP is to describe the method used by Katahdin Analytical Services technical personnel to measure the concentration of gasoline range organics in water and soil. This procedure measures compounds from MTBE through naphthalene inclusive. This corresponds to a boiling point range between approximately 60°C and 220°C. The analytical procedure is based on a purge-and-trap, Gas Chromatography (GC) procedure.

#### 1.1 Definitions

ANALYTICAL BATCH: 20 or fewer samples which are analyzed together with the same method sequence and the same lots of reagents and with the manipulations common to each sample within the same time period or in continuous sequential time periods.

METHOD BLANK (LABORATORY REAGENT BLANK): An artificial sample designed to determine if method analytes or other interferences are present in the laboratory environment, the reagents, or the apparatus. For aqueous samples, reagent water is used as a blank matrix. For soil samples, clean (muffled) sand is used as a blank matrix. The blank is taken through the appropriate steps of the process.

GASOLINE RANGE ORGANICS (GRO): All chromatographic peaks eluting from methyl-tert-butylether through naphthalene, inclusive. Quantitation is based on a direct comparison of the area within this range to the total area of the 10 components in the Gasoline Component Standard.

GASOLINE COMPONENT STANDARD: A ten component blend of typical gasoline compounds (Table 4). This standard serves as a quantitation standard and a retention time window for gasoline range organics.

INDEPENDENT CALIBRATION VERIFICATION (ICV): The ICV is obtained from a source external to the laboratory and different from the source of calibration standards. A reagent water blank is spiked with the ICV Standard and analyzed immediately following a calibration.

COMMERCIAL GASOLINE STANDARD: The Commercial Gasoline Standard is analyzed following a calibration and is used for fingerprinting purposes.

LABORATORY CONTROL SAMPLE (GASOLINE COMPONENT SPIKE): A reagent water blank or reagent methanol blank sample spiked with the Gasoline Component Standard and run through the method with samples as a quality control check. The LCS is obtained from a source external to the laboratory and different from the source of calibration standards. It is used to check laboratory performance with external prepared test materials.

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TITLE: METHOD FOR DETERMINING VOLATILE PETROLEUM HYDROCARBONS or GASOLINE RANGE ORGANICS (GRO) BY METHOD 8015 (WITH OR WITHOUT MODIFICATIONS)

MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD): An aliquot of an environmental sample to which known quantities of the method analytes are added in the laboratory. The MS/MSD is analyzed exactly like a sample, and its purpose is to determine whether the sample matrix contributes bias to the analytical results. The background concentrations of the analytes in the sample matrix must be determined in a separate aliquot and the measured values in the MS/MSD corrected for background concentrations.

METHOD DETECTION LIMIT (MDL): Minimum concentration that an analyte can be measured and reported with a 99% confidence that the analyte concentration is greater than zero and is determined from analysis of a sample in a given matrix containing the analyte. The MDL is determined using EPA Appendix B to Part 136, CFR 40 Ch. 1 (7-1-94) using the Student t Test.

PRACTICAL QUANTITATION LIMIT (PQL): The practical quantitation limit for gasoline is 10 ug/L for water samples and 2.5 mg/Kg for soil samples.

TEMPERATURE BLANK: (If required by project) A vial of water supplied by the laboratory, treated in the same manner as sample vials and carried along with samples, to determine if proper cooling of samples has been achieved (0°C-6°C). A 40 mL or 60 mL vial will be adequate for this purpose.

KATAHDIN INFORMATION MANAGEMENT SYSTEM (KIMS): A complete multi-user system with the capabilities of integrating laboratory instrumentation, generating laboratory worksheets, providing complete Lab Order status and generating reports. KIMS utilizes these features through a database.

PE NELSON TURBOCHROM OR HP CHEMSTATION: data acquisition systems that are used to collect chromatographic data. The systems can also be used to archive raw data files.

TARGET: A software system that combines full processing, reporting and comprehensive review capabilities, regardless of chromatographic vendor and data type.

TARGET DB: An oracle database used to store and organize all Target data files.

QUICKFORMS: A laboratory reporting software for Target and Target DB. The QuickForms report module for Target is preconfigured with generalized forms and US EPA CLP report forms and disk deliverables, which can be customized.

#### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of analysts experienced in

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TITLE: METHOD FOR DETERMINING VOLATILE PETROLEUM HYDROCARBONS or GASOLINE RANGE ORGANICS (GRO) BY METHOD 8015 (WITH OR WITHOUT MODIFICATIONS)

the analysis of GRO's by Method 8015C (with or without modifications). Each analyst must demonstrate and document their ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, current revision, "Personnel Training & Documentation of Capability".

It is the responsibility of all Katahdin technical personnel involved in analysis of GRO's by Method 8015 (with or without modifications) to read and understand this SOP, to adhere to the procedures outlined, and to properly document their data in the appropriate lab notebook. Any deviations from the test or irregularities with the samples should also be recorded in the lab notebook and reported to the Department Manager or designated gualified data reviewer responsible for this data.

It is the responsibility of the Department Manager to oversee that members of their group follow this SOP, to ensure that their work is properly documented and to initiate periodic review of the associated logbooks.

#### 1.3 Safety

Users of this procedure must be cognizant of inherent laboratory hazards, proper disposal procedures for contaminated materials and appropriate segregation of hazardous wastes. The toxicity or carcinogenicity of each reagent used in this method has not been precisely defined; however, each chemical should be treated as a potential health hazard. A reference file of material safety data sheets is available to all personnel involved in the chemical analysis. Everyone involved with the procedure must be familiar with the MSDSs (material safety data sheets) for all the materials used in this procedure.

Each qualified analyst or technician must be familiar with Katahdin Analytical Environmental Health and Safety Manual including the Katahdin Hazardous Waste Plan and must follow appropriate procedures. These include the use of appropriate personal protective equipment (PPE) such as safety glasses, gloves and lab coats when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location of all safety equipment. Each analyst shall receive a safety orientation from their Department Manager, or designee, appropriate for the job functions they will perform.

#### 1.4 Pollution Prevention/Waste Disposal

Whenever possible, laboratory personnel should use pollution prevention techniques to address their waste generation. Refer to the current revision of the Katahdin Hazardous Waste Management Program for further details on pollution prevention techniques.

Wastes generated during the preparation of samples must be disposed of in

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TITLE:

METHOD FOR DETERMINING VOLATILE PETROLEUM HYDROCARBONS OF GASOLINE RANGE ORGANICS (GRO) BY METHOD 8015 (WITH OR WITHOUT MODIFICATIONS)

accordance with the Katahdin Analytical Environmental Health and Safety Manual and SOP SD-903, "Sample Disposal," current revision. Expired standards are lab packed, placed in the Katahdin hazardous waste storage area, and disposed of in accordance with this SOP. Purge vial and methanol waste are disposed of in the "A" waste satellite accumulation area located between the Gasoline Range instruments.

#### 2.0 SUMMARY OF METHOD

The method provides gas chromatographic conditions for the detection of volatile petroleum fractions such as gasoline, Stoddard solvent or mineral spirits. Samples are analyzed utilizing purge-and-trap sample concentration. The gas chromatograph is temperature programmed and utilizes a capillary column (75 meter 0.45mm ID DBVRX or equivalent) to facilitate separation of organic compounds. Detection is achieved by a flame ionization detector (FID) or photoionization detector (PID)/(FID) in series with the non-destructive (PID) first in the series. Quantitation is based on FID detector response to a gasoline component standard utilizing an external standard method.

The method is suitable for the analysis of waters, soils, or wastes. Water samples can be analyzed directly for gasoline range organics by purge-and-trap extraction and gas chromatography. Soil or waste samples are dispersed in methanol to dissolve the volatile organic constituents. A portion of the methanol solution is then analyzed by purge-and-trap GC.

This method is based in part on 1) API's PETROLEUM HYDROCARBON METHODS [Revised August 1993], 2) Wisconsin GRO and DRO methods, 3) SW-846 methods 5030, 8000, and 8015; 4) The Massachusetts TPH methodologies, 5) Maine HETL GRO methods.

#### 3.0 INTERFERENCES

Heavier petroleum products such as diesel fuel may contain some volatile components, producing a response within the retention time range for GRO. Other compounds that respond to the FID such as chlorinated and oxygenated hydrocarbons are detected by this method and will be included in the concentration. If the analyst suspects that compounds are present that are not present in gasoline mixtures, the analyst should suggest additional analysis. Spills of known specific constituents should be analyzed and quantified by a method specific for those constituents.

Samples can become contaminated by diffusion of volatile organics through the sample container septum during shipment and storage or by dissolution of volatiles into the methanol used for preservation. Trip blanks prepared from both reagent water and methanol should be carried through sampling and subsequent storage and handling to serve as a check on such contamination. (Methanol trip blank required only if soil samples

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are preserved with methanol in the field.)

Contamination by carryover can occur whenever high-level and low-level samples are sequentially analyzed. To reduce carryover, the sample syringe and/or purging device must be rinsed between samples with reagent water or solvent. For volatile samples containing high concentrations of water-soluble materials, suspended solids, high boiling compounds or organohalides, it may be necessary to wash the syringe or purging device with a detergent solution, rinse with distilled water, and then dry in a 105°C oven between analyses. The trap and other parts of the system are also subject to contamination, therefore, frequent bake-out and purging of the entire system may be required. A screening step is recommended to protect analytical instrumentation. Whenever an unusually concentrated sample is encountered, it should be followed by an analysis of a water blank to check for cross-contamination.

The retention time window definition (methyl-tert-butylether through naphthalene inclusive) introduces a negative bias, however, it improves comparability between laboratory data. Note that gasoline blends often contain 10% ethanol, which could be responsible for a portion of this negative bias.

The flame ionization detector (FID) is a non-selective detector. There is a potential for many non-target compounds present in samples to interfere with this analysis. There is also the potential for analytes to be resolved poorly, especially in samples that contain many analytes. The data user should consider this and may wish to alter the target analyte list accordingly.

#### 4.0 APPARATUS AND MATERIALS

- 4.1 Gas Chromatograph Analytical system complete with gas chromatograph suitable for purge-and-trap sample introduction and all required accessories, including detectors, column supplies, recorder, gases and syringes. A data system capable of determining peak areas by integrating from baseline to baseline is required.
- 4.2 Columns (Capillary Columns are Required)
  - 4.2.1 The capillary column must be capable of resolving typical gasoline components. It must be capable of achieving a 60% resolution of all 10 components, with the exception of meta and para-xylene. It must also be capable of separating methyl-tert-butylether from methanol at the concentration resulting from preparation of the standard or in the sample spiked with surrogate. The Resolution is defined as:

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- 4.2.2 Suggested columns: Note either column can be used for soil or water provided the resolution meets the criteria given above.
  - DBVRX 75 meter 0.45 mm ID, 2.5 um film thickness (or equivalent).
  - RTX 502.2 105 meter 0.53 mm ID, 3.0 um film thickness (or equivalent).
  - Any capillary column phase and length and diameter may be used as long as the requirements of resolution of this method are met.
- 4.3 Detector: Flame ionization (FID), or FID in series with a Photoionization detector (PID).
- 4.4 Purge-and-trap device: The purge-and-trap device consists of three separate pieces of equipment: the sample purger, the trap, and the desorber. Several complete devices are commercially available.
  - 4.4.1 Commercially available automated sample purge devices may be used, provided that equivalent performance is demonstrated.
  - 4.4.2 The recommended trap is the Supelco H trap which consists of 7.6 cm Carbopack B and 1.3 cm Carbosieve S-III. Refer to the instrument runlog for the trap conditions. The trap length and packing materials may be varied as long as equivalent performance compared to the recommended trap has been verified. Demonstration of equivalency of trap performance must be based on a full range of gasoline components, not just on the 10 components in the standard.
  - 4.4.3 Traps should be conditioned, desorbed, and baked according to the manufacturers' guidelines. The trap may be vented to the analytical column during daily conditioning, however, the column must be run through the temperature program prior to analysis of samples.
  - 4.4.4 The desorber should be capable of rapidly heating the trap to the recommended desorption temperature.
- 4.5 Analytical balance: A balance capable of accurately weighing 0.0001 g (for preparing standards). A top-loading balance capable of weighing to the nearest 0.01 g (for weighing soil samples).
- 4.6 Ultrasonic bath.
- 4.7 VOC Vials: 40 mL VOC vials with Teflon/silicone septa for waters and soils. Soils

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not preserved in the field may be collected in wide mouth jars with Teflon lined caps (2 or 4 oz jars recommended).

- 4.8 Syringes: 5 mL gas-tight syringe (with shutoff valve recommended).
- 4.9 Syringe valve: Two-way, with luer ends.
- 4.10 Volumetric flask (class A): 10 mL, 50 mL, 100 mL, 500 mL, and 1,000 mL with a ground-glass or screw-top stopper.
- 4.11 Microsyringes: 1 uL, 5 uL, 10 uL, 25 uL, 100 uL, 250 uL, 500 uL, and 1,000 uL.
- 4.12 Disposable pipettes: Pasteur.
- 4.13 Spatula: Stainless steel.
- 4.14 40 mL VOA vials with Teflon lined septa or caps for soil extractions performed in the lab.

#### 5.0 REAGENTS

- 5.1 Reagent Water: Organic free water.
- 5.2 Methanol: Purge and trap grade or equivalent. Store away from other solvents.
- 5.3 Acid for preserving water samples: A 1:1 mixture of reagent water and concentrated hydrochloric acid. Use 2 or more drops per 40 mL VOA vial. Acid may be added to the sample at the time of collection or may be added to the vial prior to the collection. Alternatively add 0.1 g of sodium hydrogen sulfate to the empty VOA vial. The final pH of the water should be <2.
- 5.4 GRO free sand or soil.
- 5.5 Stock Standards: Purchase individual certified component standards. A concentration of 20 mg/mL is recommended for individual component standards.
  - 5.5.1 Transfer the stock standard solution into a Teflon-sealed screw-cap or crimp cap bottle. Refrigerate, with minimal headspace and protect from light.
  - 5.5.2 Standards must be replaced after 6 months or sooner if comparison with check standards indicates a problem.
- 5.6 Gasoline Component Standard: Purchase a certified Gasoline Component Standard

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from an acceptable supplier like Supelco. These standards should be stored with minimal headspace and should be checked frequently for signs of degradation or evaporation. The component standard must be replaced after 6 months or sooner if comparison with a check standard indicates a problem.

- 5.7 Calibration Standards: Prepare Calibration standards at a minimum of five concentration levels in reagent water from the Gasoline Component Standard. One of the concentration levels should be at the minimum reporting level. The remaining concentration levels should correspond to the working range of the GC system. Refer to section 7.4.
- 5.8 Independent Calibration Standard (ICV)/Laboratory Control Spike Standard (LCS) Stock Standards: Purchased form a source external to the laboratory and different from the source of calibration standards. These standards should be stored with minimal headspace and should be checked frequently for signs of degradation or evaporation. The component standard must be replaced after 6 months or sooner if comparison with a check standard indicates a problem.
- 5.9 Independent Calibration Standard (ICV)/Laboratory Control Spike Standard (LCS) Working Standards: Prepare standards at a concentration level near or at the midpoint of the calibration in methanol from the Independent Calibration Standard (ICV)/Laboratory Control Spike Standard (LCS) Stock Standards.
- 5.10 Surrogate Control Standard (SCS): The analyst should monitor the performance of the analytical system by spiking each water sample, standard, water blank and diluted soil extract with the surrogate compound Bromofluorobenzene (BFB). All samples are spiked with 2 uL of a 50 ug/mL concentration of BFB to give a final concentration of 20 ug/L.
- 5.11 Commercial Gasoline: Purchase a certified Commercial Gasoline Fuel Standard from an acceptable supplier like Supelco. This standard should be stored with minimal headspace and should be checked frequently for signs of degradation or evaporation. The standard must be replaced after 6 months or sooner if comparison with a check standard indicates a problem.

#### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

- 6.1 Soil core samples may be collected in wide mouth jars when field preservation is not required. Minimum handling is required to reduce VOC loss. Samples that are preserved in the field should be collected in septum vials.
  - 6.1.1 Note on the chain of custody when samples are not preserved in the field. If the sample is not preserved in the field, the container should be filled to

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minimize the headspace. If there is a large headspace, this fact should also be noted on the chain of custody.

- 6.1.2 If samples are not received cold, note this occurrence on the chain of custody.
- 6.1.3 Refrigerate samples at (4°C ± 2°C) as soon as possible after they are received at the lab.
- 6.1.4 Analysis must be completed within 14 days of collection.
- 6.2 Collect water samples in 40 mL septum vials and preserve with acid at time of collection.
  - 6.2.1 Check water samples for air bubble(s) and reject if the bubble(s) are significant.
  - 6.2.2 If samples are not received cold, note this occurrence on the chain of custody.
  - 6.2.3 Place samples in a refrigerator at 4°C ± 2°C as soon as possible after they are received at the lab.
  - 6.2.4 A water trip blank that accompanies all samples and is analyzed with the samples may be required for specific projects.
  - 6.2.5 Water samples must be analyzed within 14 days of collection.

#### 7.0 PROCEDURES

7.1 Volatile compounds are introduced into the gas chromatograph by purge-and-trap. Purge-and-trap may be used directly on groundwater samples. Soils and solids should be analyzed by methanol extraction followed by purge-and-trap. Soil concentrations will be reported on a dry weight basis.

#### 7.2 Gas Chromatography

7.2.1 Conditions for Column 1: Suggested conditions for a 75 m x 0.45 mm I.D. DBVRX, or equivalent: Set helium column pressure as recommended by the manufacturer. Set column temperature to 35°C for 7 min, then 4°C/min to 130°C, then 8°C/min to 160°C, 20°C/min to 230°C (hold for 4 min). Conditions may be altered to improve resolution of gasoline range organics.

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7.2.2 Other columns: Set GC conditions to meet the criteria in Section 4.2.1.

- 7.3 Retention Time Window and Quantitation
  - 7.3.1 Gasoline Range Organics (GRO): All chromatographic peaks eluting from methyl-tert-butylether through naphthalene, inclusive. Quantitation is based on a direct comparison of the area within this range to the total area of the 10 components in the Gasoline Component Standard. (Using a "baseline to baseline" integration as opposed to a "valley to valley" integration.)
  - **Note:** The area of the MTBE peak may be determined by tangent skimming when necessary.
  - 7.3.2 The retention time window is defined as beginning approximately 0.1 minutes before the retention time of methyl-tert-butylether and ending 0.1 minutes after the retention time of naphthalene in the calibration run.

Please note that using the nominal window of 0.1 minutes and the use of the retention time markers methyl-tert-butylether and naphthalene may not be allowable for certain states, federal programs, or clients. For projects requiring DoD QSM, current version,, the retention time width is plus or minus three times the standard deviation for each analyte retention time from a 72-hour study. South Carolina only allows the use of established retention time windows and the retention time markers 2-Methylpentane and 1,2,4-Trimethylbenzene. The retention time windows must be established as described below.

- 7.3.2.1 Three injections of the GRO component standard mix are made throughout the course of a 72 hour period.
- 7.3.2.2 The standard deviation of the three retention times for 2-Methylpentane and 1,2,4-Trimethylbenzene are calculated.
- 7.3.2.3 Plus or minus three times the standard deviation of the retention times for each standard is used to define the retention time window.
- 7.3.2.4 Retention time windows are calculated for each standard on each GC column and whenever a new GC column is installed. The data is kept on file in the laboratory.
- 7.3.2.5 If the calculated retention time window results in a value of 0.03 minutes or less, the laboratory will apply a nominal window of 0.03 minutes.
- 7.3.3 The laboratory must determine retention time windows for the first and last

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standard on each GC column and whenever a new GC column is installed. This data must be retained by the laboratory.

7.3.4 Quantify by summing all peak areas eluting from methyl-tert-butylether through naphthalene, inclusive.

#### 7.4 GRO Calibration

7.4.1 Run the Gasoline Component Standard at a minimum of five concentration levels at the minimum reporting level and covering the working range of the instrument. When the calibration curve is run an independent check standard should also be run to validate the curve. A gasoline standard is also run with the calibration curve for fingerprinting purposes.

Please note that using the Gasoline component calibration standard may not be allowable for certain states, federal programs, or clients. South Carolina requires the use of a fuel oil as the calibration standard. If the specific fuel type is known, the laboratory should obtain a pure sample of material from the tank that is leaking. For unknown samples, the instrument is calibrated using Gasoline fuel. The calibration levels for Gasoline are prepared at the same concentrations as that of the GRO calibration. The 100 ug/L standard is used as the mid-point or calibration verification standard (CV).

7.4.2 The instrument is calibrated by injecting a specific amount of one of the three mixes as indicated in the table below. Mix A contains the GRO component mix at 25 ug/mL and the surrogate BFB at 5 ug/mL. Mix B contains the GRO component mix at 250 ug/mL and the surrogate BFB at 50 ug/mL. Mix C contains the GRO component mix at 2500 ug/mL and the surrogate BFB at 500 ug/mL.

Amount injected	Concentration of GRO ug/L	Concentration of BFB ug/L
2 uL of Mix A	10	2
2 uL of Mix B	100	20
5 uL of Mix B	250	50
10 uL of Mix B	500	100
2 uL of Mix C	1000	200
4 uL of Mix C	2000	400

The solutions are added directly in the 5 mL glass syringe as follows:

 Add the aliquot of calibration solution directly to the reagent water in the glass syringe by inserting the needle through the syringe end.

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- When discharging the contents of the microsyringe, be sure that the end of the syringe needle is well beneath the surface of the reagent water.
- Inject the standard into the purge vessel through the two-way valve.
- 7.4.3 Inject 5 mLs of each calibration standard utilizing the purge-and-trap analysis. Retention time window position shall be set using the midpoint standard from the curve. Tabulate the entire peak area (baseline to baseline) for the ten components against the mass injected. The results are used to prepare a calibration curve by linear regression.
- 7.4.4 An Independent Calibration Verification Standard is analyzed immediately after calibration, before any samples are analyzed. For projects requiring DoD QSM, current version,, all project analytes must fall within the established retention time windows.
- 7.4.5 A gasoline standard is analyzed following the calibration curve to be used for fingerprinting purposes.
- 7.4.6 The working calibration curve must be verified at the beginning and end of each analytical sequence by the injection of a mid-point Gasoline Component Standard or calibration verification standard (CV). A CV must also be analyzed every 10 samples or after a 12 hour shift, whichever is sooner. If the response for the CV varies from the predicted response by more than 20%, the analytical system should be examined to determine the cause and corrective action should be performed and/or a new CV should be prepared and analyzed. If the 20% criteria is still not met, the system must be recalibrated. All samples must be reanalyzed that fall within the standard that exceeded criteria and the last standard that was acceptable.

  The closing CV should be at a level different than the opening CV.

Please note that using the Gasoline component calibration verification standard (CV) as well as an acceptance criteria of 20% may not be allowable for certain states, federal programs, or clients. For projects requiring DoD QSM, current version,, all project analytes must fall within the established retention time windows. South Carolina requires the use of a fuel oil as the CV. Also, South Carolina requires the analysis of a standard that contains 2-Methylpentane and 1,2,4-Trimethylbenzene at the beginning and end of each 12 hour work shift. The acceptance criteria for the fuel oil CV is 15% and 2-Methylpentane and 1,2,4-Trimethylbenzene must fall within the established retention time window.

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7.5 Gas Chromatographic Analysis of Water Samples

Please refer to Katahdin Analytical Services SOP CA-108, "Basic Laboratory Technique", current revision, for more information on subsampling.

- 7.5.1 Introduce volatile compounds into the gas chromatograph using the purgeand-trap method. For automated systems follow the manufacturers' recommended procedure.
- 7.5.2 Adjust the purge gas flow rate (nitrogen or helium) to 25-40 mL/min on the purge-and-trap device.
- 7.5.3 Remove the plunger from a 5 mL gas-tight syringe that has been rinsed three times with reagent water. Open the sample or standard bottle and carefully pour the sample into the syringe. Replace the plunger and vent any residual air while adjusting the sample volume to 5 mL. Care must be taken to prevent air from leaking into the syringe.
- 7.5.4 For samples requiring dilutions, the sample may be diluted directly in the 5 mL gas-tight syringe. Alternatively, the following procedure is appropriate for diluting purgeable samples. All steps must be performed without delays until the diluted sample is in a gas-tight syringe.
- 7.5.5 Dilutions may be made in volumetric flasks (10 mL to 100 mL). Select the volumetric flask that will allow for the necessary dilution. Intermediate dilutions may be necessary for highly concentrated samples.
- 7.5.6 Calculate the approximate volume of reagent water to be added to the volumetric flask selected and add slightly less than this volume of reagent water to the flask.
- 7.5.7 Inject the proper aliquot of samples (taken from a filled VOA vial or from a storage syringe prepared as in Section 7.5.3) into the flask. If aliquots of less than 1 mL are required, use microliter syringes and deep injections to transfer the sample aliquot to the flask. Dilute the sample to the mark with reagent water. Cap the flask and invert three times. Repeat the above procedure for additional dilutions. Alternatively the dilutions can be made directly in a gas tight syringe to avoid further loss of volatiles.
- 7.5.8 Fill a 5 mL syringe with diluted sample as in Section 7.5.3.
- 7.5.9 For aqueous LCSs, use a 5 mL syringe rinsed three times with reagent water. The volume is then brought to 5 mL and 2 uL of a 50 ug/mL concentration of BFB and 2 uL of a 250 ug/mL concentration of the GRO

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LCS Mix are added to the 5 mL syringe. For the MS and/or MSD, after rinsing with water, the syringe is rinsed with sample and then is brought to 5 mL with sample. The same amount of the same standards used for the LCS are added to the 5 mL syringe of sample.

- 7.5.10 Add surrogate to sample or diluted sample by spiking 2 uL of a 50 ug/mL solution of BFB directly into the syringe. Inject sample into the purging chamber.
- 7.5.11 Close the valve and purge the sample for 11 minutes at ambient temperature.
- 7.5.12 At the conclusion of the purge time, the purge and trap concentrator should wait in desorb ready mode until the GC temperature stabilizes at 35°C. The purge-and-trap will then be triggered by the GC as it initiates its run and the concentrator will move automatically into the desorb mode. As desorption is initiated the trapped materials are introduced to the gas chromatographic column by rapidly heating the trap to the manufacturer's recommended desorption temperature and backflushing the trap for the recommended desorption time.
- 7.5.13 Wash the ALS position with a minimum of one 5 mL flush of reagent water (or methanol followed by reagent water) to avoid carryover of pollutant compounds into subsequent analyses.
- 7.5.14 After desorbing the sample, recondition the trap by baking the purge-and-trap device at the recommended bake temperature. After approximately 7-35 min, turn off the trap heater and open the syringe valve to stop the gas flow through the trap. When cool, the trap is ready for the next sample.
- 7.5.15 If the initial analysis of a sample or a dilution of the sample has a concentration of analytes that exceeds the initial calibration range, the sample must be reanalyzed at a higher dilution. When a sample is analyzed that has a saturated response from a compound, this analysis must be followed by a blank reagent water analysis. If the blank analysis is not free of interferences, the system must be decontaminated. Sample analysis may not resume until a blank can be analyzed that is free of interferences.
- 7.5.16 All dilutions should keep the response of the major constituents (previously saturated peaks) in the linear range of the curve.
- 7.5.17 All water samples should be checked to make sure that they were preserved. After sample analysis (or after removing an aliquot for analysis), check the pH of the water using pH paper. If the pH is not <2, this fact

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should be noted in the instrument run-log, and noted in the sample narrative.

#### 7.6 Methanol Extraction for Soil/Sediment

- 7.6.1 This method is based on extracting the sediment/soil with methanol. An aliquot of the extract is added to reagent water and purged at the conditions indicated in Table 3.
  - 7.6.1.1 Add approximately 10 g of sample into a tared sample vial and weigh to the nearest 0.01 g. Immediately add 10 mLs of methanol to the vial.
  - 7.6.1.2 If the soil sample has been methanol preserved in the field, both the weight of the soil and the volume of the methanol must be recorded. The ratio of the soil to methanol should be approximately 1 gram soil: 1 milliliter of methanol with a minimum soil weight of 10 grams.
  - 7.6.1.3 For the LCS/LCSD, 10 g of sand is added to a 40 mL VOA vial followed by the addition of 10 mL of methanol. Then 12.5 uL of a 2000 ug/mL concentration of Gasoline Component Spike solution is spiked into the methanol.
  - 7.6.1.4 The MS/MSD is prepared by adding 10 g of sample into a 40 mL VOA vial followed by 10 mL of methanol. Then 12.5 uL of a 2000 ug/mL concentration of Gasoline Component Spike solution is spiked into the methanol.
  - 7.6.1.5 Record all weights and volumes in the GC Soil Prep Logbook (Figure 1).
- 7.6.2 Shake the sample(s) and QC for 2 minutes and sonicate for 20 minutes.
- 7.6.3 Allow sediment to settle until a layer of methanol is apparent. Centrifuge if necessary. If not analyzed immediately store at 6°C or less.
- 7.6.4 Using a microliter syringe, withdraw an appropriate volume.
- 7.6.5 Remove the plunger from a 5 mL gas tight syringe equipped with a syringe valve that has been rinsed three times with reagent water and fill until overflowing with reagent water. Replace the plunger and compress the water to vent trapped air. Pull the plunger to 5 mLs and add 20 uL of the sample extract.

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- 7.6.6 Add surrogate to sample or diluted sample by spiking 2 uL of a 50 ug/mL solution of BFB directly into the syringe. Inject sample into the purging chamber.
- 7.6.7 Proceed with the analysis as in 7.5.11-7.5.16. Analyze all reagent blanks and QC samples on the same instrument as that used for the samples. The reagent blank should contain 20 uL of the methanol used to extract the samples (or a volume equal to the largest amount of extract purged).
- 7.6.8 If the responses exceed the calibration or linear range of the systems, repeat the analysis using a smaller aliquot of methanol extract.

#### 7.7 Calculations

7.7.1 GRO Calibration: The concentration of Gasoline Range Organics in the sample is determined from a summation of the total peak area for all chromatographic peaks eluting from methyl-tert-butylether through naphthalene, inclusive, using the calibration curve. (Using a "baseline to baseline" integration as opposed to a "valley to valley" integration.) Refer to Section 7.3 (Retention Time Windows and Quantitation).

Quantitation may be based on a linear regression equation derived from the calibration curve.

Linear regression: From the calibration standards GC responses, R, and their known concentrations, C in ug/L, the following linear equation may be derived [R is plotted on the y axis; C is plotted on the x axis]:

R = mC + b; which can be rearranged to C = (R - b)/(m).

Using the slope (m) and the intercept (b) from this equation the concentration of the sample can be calculated.

- 7.7.2 The concentration of an analyte is calculated by using the calibrated curve that is prepared in Target. When an analyte is identified, Target displays a concentration when the file is processed through the appropriate calibrated method.
- 7.7.3 The concentrations from the reports are then incorporated to arrive at a final sample concentration.

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Water: Concentration (ug/L) = (C) (5mL)/(Vs)

Soil: Concentration (mg/Kgdrywt) = (C)  $(V_m)(0.005L)/(Ws)(V_E)(D)$ 

Where: C = Concentration calculated by Turbochrom in ug/L

Vs = Volume of sample purged in mL Ws = Weight of sample extracted in grams

V<sub>M</sub> = Volume of methanol in mLV<sub>E</sub> = Volume of extract purged in mL

D = Decimal total solids

- 7.7.4 Blank areas may not be subtracted from sample areas. Chromatographic baseline rises due to temperature programming may be corrected for by using baseline correction. The baseline correction may be performed by the most convenient method that the data handling system allows.
- 7.7.5 If an aqueous method blank exceeds the PQL of 10 ug/L, all water samples associated with this blank must be rerun. However, if the sample concentration exceeds the blank contamination by 10 times, the contribution from the blank is negligible, and the data may be reported with narration. For projects requiring DoD QSM, current version,, the method blank must not detect any analytes >1/2 the PQL and >1/10 the amount measured in any sample or 1/10 the regulatory limit (whichever is greater).
- 7.7.6 If a methanol blank exceeds the PQL of 2.5 mg/Kg, all soil samples associated with this blank must be rerun. However, if the sample concentration exceeds the blank contamination by 10 times, the contribution from the blank is negligible, and the data may be reported with narration. For projects requiring DoD QSM, current version,, the method blank must not detect any analytes >1/2 the PQL and >1/10 the amount measured in any sample or 1/10 the regulatory limit (whichever is greater).
- 7.7.7 To ensure that peaks outside the GRO window are not missed, run the chromatogram out 5 minutes past the last component in the GRO component standard. All significant peaks (and baseline rises) outside the window should be qualitatively assessed.

#### 7.8 Data Review

#### 7.8.1 Initial Data Review

7.8.1.1 The initial data review is accomplished by the analyst who ran the samples. This review is of sufficient quality and detail to provide a list of samples that need to be reanalyzed or diluted and reanalyzed.

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The initial data review is performed on the detailed quantitation reports of the analyzed samples. This data review examines criteria that directly impact whether or not the sample needs to be reanalyzed and/or reextracted. These criteria include:

- QC criteria for method blank, LCS, MS/MSD, and calibration refer to section 8.0.
- Surrogate recovery.
- Chromatography: manual integration.
- Target compound detection: quantitation, false positives.
- 7.8.1.2 The requirement of the GC laboratory is that this initial data review be completed no later than the end of the next work day. After the analyst has completed his or her initial data review, the information is then ready to be processed for reporting. Refer to section 7.9.

#### 7.8.2 Surrogate recovery

- 7.8.2.1 All recoveries must meet the most recent laboratory established acceptance limits, which are listed on the GC Laboratory Surrogate Acceptance Limit sheet.
- 7.8.2.2 The sample is evaluated for recovery of the surrogate BFB for water samples and soil samples. For both water and soil samples, if the recovery of BFB is low and there is no apparent matrix effect, the sample should be reanalyzed. If the BFB is low and there may be a matrix effect, reanalyze to confirm a matrix effect. If the surrogate is high and the sample results are less than the PQL, or there is likely a matrix effect, narrate.
- 7.8.2.3 For projects requiring DoD QSM, current version,, Q-flag all detected analytes in the sample if the surrogates fail the acceptance criteria.

#### 7.8.3 Chromatography

- 7.8.3.1 The chromatography should be examined for the presence of any non-target peaks, which can be used as an indication of whether or not matrix interference might be influencing surrogate recoveries.
- 7.8.3.2 Manual integrations are to be performed when chromatographic conditions preclude the computer algorithm from correctly integrating

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the peak of concern. In no instance shall a manual integration be performed solely to bring a peak within criteria.

7.8.3.3 Each peak of concern is examined by the primary analyst to ensure that the peak was integrated properly by the computer algorithm. Any manual integrations that are necessary (for instance, if the sample contains a concentration of GRO which was integrated "valley to valley" instead of a "baseline to baseline"), are performed in Target Review. An "M" qualifier will automatically be printed on the quantitation report summary. For specific procedures on how to manually integrate, refer to Katahdin SOP QA-812, Manual Integration, current revision.

#### 7.8.4 Target Compound Detection

To correctly quantitate the GRO concentration, the area of the surrogate BFB is not included in the GRO timed range area. In some instances, BFB may need to be manually integrated in Target Review to correctly quantitate the BFB and GRO concentrations.

#### 7.9 Reporting

After the chromatograms have been reviewed and any target analytes have been quantitated using Target, the necessary files are brought into QuickForms. Depending on the QC level requested by the client, a Report of Analysis (ROA) and additional reports, such as LCS forms and chronology forms, are generated. The package is assembled to include the necessary forms and raw data. The data package is reviewed by the primary analyst and then forwarded to the secondary reviewer. The secondary reviewer validates the data and checks the package for any errors. When completed, the package is sent to the department manager for final review. A completed review checklist is provided with each package. The final data package from the Organics department is then processed by the Data Management department.

#### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

Refer to Table 1 for a summary of QC requirements, acceptance criteria, and corrective actions. Table 1 criteria are intended to be guidelines for analysts. The table does not cover all possible situations. If any of the QC requirements are outside the recovery ranges listed in Table 1, all associated samples must be evaluated against all the QC. In some cases data may be reported, but may be reanalyzed in other cases. Making new reagents and standards may be necessary if the standardization is suspect. The corrective actions listed in Table 1 may rely on analyst experience to make sound scientific judgments. These decisions are

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based on holding time considerations, client and project specific Data Quality Objectives and on review of chromatograms. The Department Manager, Operations Manager, and/or Quality Assurance Officer may be consulted to evaluate data. Due to time constraints, samples may not be able to be reanalyzed within hold time. In these cases "qualified" data with narration may be advisable after consultation with the client.

In some cases the standard QC requirements listed in this section and in Table 1 may not be sufficient to meet the Data Quality Objectives of the specific project. Much of the work performed at the lab is analyzed in accordance with specific QC requirements spelled out in a project specific Quality Assurance Project Plan (QAPP) or in a program specific Quality Systems Manual (QSM). The reporting limits, acceptance criteria and/or corrective actions may be different than those specified in this SOP. In these cases the appropriate information will be communicated to the Department Manager and/or senior chemists before initiation of the analyses so that specific product codes can be produced for the project. In addition, the work order notes for each project will describe the specific QAPP or QSM to be followed.

- 8.1 The analyst must make an initial demonstration of the ability to generate acceptable accuracy and precision with this method by successful analysis of the following:
  - 8.1.1 Replicate Commercial Gasoline Spikes in Water: Analysis of 4 replicates at a concentration of 100 ug/L (in water) with an accuracy falling between 60% and 140% of the known concentration with a precision of 20% or less.
  - 8.1.2 Replicate Commercial Gasoline Spikes in Soil: Analysis of 4 replicates at a concentration of 25 mg/Kg with an accuracy between 60% and 140% of the known concentration and the precision should be within 20%. Soil spikes should be prepared and analyzed as described in Section 7.6.1.3.
- 8.2 For every 20 samples analyzed the lab must analyze a set of duplicate Gasoline component Spikes in water. The duplicate spikes must be run through the method in the same manner as samples. The accuracy of the two water spikes must be within the laboratory's statistically derived acceptance limits. and the percent relative difference between the two values must be 20% or less.
  - Projects requesting DoD QSM, Current Version, require Q-flagging the specific LCS analytes that fail and are detected in the associated samples. MS/MSD failures require a J-flag in the parent sample for the analytes that fail the acceptance criteria.
- 8.3 With every analytical sequence containing soil samples, the lab must analyze one Gasoline Component Spike in clean sand or soil. The spike amount should fall in the linear range of the detector. The spike recovery must be within the laboratory's statistically derived acceptance limits. If soils are extracted in the lab, the soil spike should be prepared at the time of the extraction and analyzed along with the samples.

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Projects requesting DoD QSM, Current Version, require Q-flagging the specific LCS analytes that fail and are detected in the associated samples. MS/MSD failures require a J-flag in the parent sample for the analytes that fail the acceptance criteria.

- 8.4 A water blank (containing purge-and-trap surrogate) must be run with every analytical sequence containing water samples, utilizing the same water used to prepare standards and make dilutions. The amount of material in the blank should not be at or above the PQL. For projects requiring DoD QSM, Current Version,, the method blank must not detect any analytes >1/2 the PQL and >1/10 the amount measured in any sample or 1/10 the regulatory limit (whichever is greater).
- A reagent methanol blank (containing purge-and-trap surrogate) prepared with clean (muffled) sand must be run with each analytical sequence containing soil samples, utilizing the same methanol used to prepare standards and dilute extracts and a methanol spike of 20 uL (or largest amount of extract used). The amount of material in the blank should not be at or above the PQL. For projects requiring DoD QSM, currnt version,, the method blank must not detect any analytes >1/2 the PQL and >1/10 the amount measured in any sample or 1/10 the regulatory limit (whichever is greater).

#### 8.6 Calibration requirements

- 8.6.1 When linear regression analysis is used for calculations the correlation coefficient (r) must be at least 0.995. For linear models, Target reports r<sup>2</sup>. This is calculated by either calculating r or squaring the result or by calculating the coefficient of determination. For a linear calibration, the equation for either is the same. The value for r<sup>2</sup> must greater than or equal to 0.990.
- 8.6.2 When average RF is used for calculations, the relative standard deviation of the RF's must not exceed 20%.
- 8.6.3 An Independent Calibration Verification Standard (ICV) (obtained from a source independent of the calibration standards) should be analyzed concurrent with the calibration standards in order to confirm the validity of the calibration curve. The ICV should fall within 20% of the expected value using the calibration data.
- 8.6.4 The calibration curve must be verified with each analytical sequence by running an opening and closing Calibration Verification Standard (CV). The closing CV is at 250 ppb. A CV must also be analyzed every 10 samples or after a 12 hour shift, whichever is sooner. The response must fall within 20% of the expected response.
- 8.7 If any of the criteria above are not met, the problem must be corrected before further

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samples are analyzed. Any samples that were analyzed following standards that did not meet calibration criteria must be reanalyzed (if reanalysis is not possible, the data must be flagged.)

8.8 Laboratory Spiked Duplicates are to be run at a frequency of 10%.

Alternatively duplicate samples and spiked samples can be substituted for the laboratory spiked duplicates at a frequency of 10%. Care must be taken to ensure that the samples are homogeneous before analyzing duplicates and spikes.

- 8.9 One methanol field blank must accompany each sampling event (for each site and each day that samples are collected and preserved in the field).
- 8.10 Trip blanks, field blanks, field duplicates and/or matrix spikes may be required for specific sampling programs.
- 8.11 Water blanks should be run after samples suspected of being highly concentrated to prevent carryover.
- 8.12 It is recommended that an acceptance criteria be established for recoveries of surrogates. Collect recoveries from 30 samples where no interference is suspected and calculate the mean recovery (X) and standard deviation (S). The acceptance limits for samples not exhibiting matrix interference will be X-3S to X+3S. The warning limits will be X-2S to X+2S. Plotting the surrogate recoveries on a control chart will make checking recoveries easier and is highly recommended. Refer to the GC Laboratory Surrogate Acceptance Limits sheet for the current limits used.

If surrogate recovery is outside of the established limits, verify calculations, dilutions, and standard solutions. Verify instrument performance, including checking for leaks and purge problems if the recovery is low. Low recovery may be due to the sample matrix. The analysis should be repeated, if the recovery is less than 50% or, if the analyses cannot be repeated, the data should be flagged.

High recoveries may be due to coeluting matrix interference. Surrogate recoveries may be reported as "masked" in high level samples exhibiting matrix interference. These samples do not need to be rerun solely to try to bring surrogate recovery into acceptance limits.

For projects requiring DoD QSM, current version,, Q-flag all detected analytes in the sample if the surrogates fail the acceptance criteria.

8.13 Non-conformance Report (NCR): Whenever data is not acceptable because of a failing LCS or surrogate recovery, a non-conformance report must be initiated as soon as possible.

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#### 9.0 METHOD PERFORMANCE

The method detection limit (MDL) is defined as the minimum concentration of a substance that can be measured and reported with 99% confidence that the value is above zero. The MDLs shall be determined and verified one time per type of instrument unless otherwise required by the method.

Limit of Detection (LOD) is an estimate of the minimum amount of a substance that an analytical process can reliably detect. An LOD is analyte and matrix specific and may be laboratory-dependent. LODs must be determined for all parameters for which the laboratory is accredited under the DoD Environmental Laboratory Accreditation Program. LOD's must be verified for every preparation and analytical method combination and on every applicable instrument on a quarterly basis.

The Limit of Detection (LOQ) is the minimum levels, concentrations, or quantities of a target variable (e.g., target analyte) that can be reported with a specified degree of confidence. The LOQ shall be set at the lowest point in the calibration curve for all analyses utilizing an initial calibration. LOQ's must be verified quarterly for every preparation and analytical method combination and on every applicable instrument on a quarterly basis for all parameters included in the DoD Scope of Accreditation. The LOQ must be verified at least once annually if the analysis is not included in the DoD Scope of Accreditation.

MDLs are filed with the Organic Department Manager and then with the QAO. LOD and LOQ verifications are filed with the QAO

Refer to the current revision of Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, for procedures on determining the MDL.

Refer to the current revision of Method 8015 for other method performance parameters and requirements.

#### 10.0 APPLICABLE DOCUMENTS/REFERENCES

"Test Methods for the Evaluation of Solid Waste: Physical/Chemical Methods", SW-846, third Edition, Final Update IV, February 2007, Method 8000B, 8015C, and 5030B.

Wisconsin DNR Modified GRO, July 1993

American Petroleum Institute, "Method for Determination of Gasoline Range Organics", 9/93

Maine HETL, Method 4.2.17, Modified Method for the Determination of GROs, 9/95.

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Katahdin SOP CA-101, Equipment Maintenance and Troubleshooting, current revision.

Department of Defense Quality Systems Manual for Environmental Laboratories (DoD QSM), current version.

The National Environmental Laboratory Accreditation Conference (NELAC) Standards, June 2003.

The NELAC Institute, Laboratory Accreditation Standards, Volume 1, Management and Technical Requirements for Laboratories Performing Environmental Analysis, 10/06/2010.

#### LIST OF TABLES AND FIGURES

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- Table 3 Purge-and-Trap Operating Parameters
- Table 4 Gasoline Standard Components and Concentrations
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- Figure 1 GC Soil Prep Logbook
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- Figure 3 Data Review Checklist

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### TABLE 1

#### QC REQUIREMENTS

Parameter/ Method	QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action
GRO by Method 8015	Method blank	One per twenty samples or every day whichever is more frequent.	GRO not detected  ≥PQL  For projects requiring  DoD QSM, current  version,, the method  blank must not detect  any analytes >1/2 the  PQL and >1/10 the  amount measured in  any sample or 1/10  the regulatory limit  (whichever is  greater).	(1) Investigate source of contamination (2) Evaluate the samples and associated QC: i.e. If the blank results are above the PQL, report sample results which are < PQL or > 10X the blank concentration. Otherwise, reprep a blank and the remaining samples.
	LCS/LCSD	One set per twenty samples or every day whichever is more frequent.	Laboratory established acceptance limits	(1) Evaluate the samples and associated QC: i.e. If an MS/MSD was analyzed and acceptable, narrate. If an LCS/LCSD was performed and only one of the set was unacceptable, narrate. If the surrogate recoveries in the LCS are low but are acceptable in the blank and samples, narrate. If the LCS recovery is high but the sample results are < PQL, narrate. Otherwise, reprep a blank and the remaining samples.
	ICAL consists of a minimum of 5 calibration standards. The low standard is at or near the PQL.	Once prior to initiating method then as needed (when CV fails).	Correlation coefficient $(r) \ge 0.995$ , $(r^2) \ge 0.990$ . See sect. 8.6.1 for further information.	(1) Perform instrument maintenance as needed and reprep and/or reanalyze the 5 calibration standards.
	CV at or near the mid-point of the calibration curve.	Beginning of each analytical sequence if ICAL previously run; end of analytical sequence; every 10 samples or after 12 hour shift, whichever is sooner	+/- 20% D	(1) Evaluate the samples: If the %D>±20% and sample results are <pql, %d="" (2)="" if="" narrate.="">±20% at the end of a sequence and is likely a result of matrix interference (i.e. samples analyzed before it had bad matrices), evaluate the samples between the opening and closing CV and narrate as needed. All samples must be reanalyzed that fall within the standard that exceeded criteria and the last standard that was acceptable.</pql,>

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#### TABLE 1, cont'd

#### **QC REQUIREMENTS**

Parameter/ Method	QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	
GRO by Method 8015	Matrix Spike/Matrix Spike Duplicate	One set for every twenty samples For projects requiring DoD QSM, current version,, one set per preparatory batch, per matrix	Laboratory established acceptance limits RPD <30	(1) Evaluate the samples and associated QC: i.e. If the LCS results are acceptable, narrate. (2) If both the LCS and MS/MSD are unacceptable, reprep the samples and QC.	
	Sample Duplicate (If required in lieu of MSD)	One sample duplicate per twenty samples	RPD <u>&lt;</u> 20	(1) If lab QC in criteria and matrix interference suspected, flag data (2) Else, reanalyze	
	Demonstration of analyst proficiency; accuracy and precision	One time per analyst initially and annually thereafter	Must pass all applicable QC for method	Repeat analysis until able to perform passing QC; document successful performance in personal training file	
	MDL study	Refer to KAS SOP QA-806, "Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications", current revision.			

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# TABLE 2 SUMMARY OF METHOD MODIFICATIONS

Topic	Katahdin SOP CA-316-10	Method 8015, current revision
Apparatus/Materials		
Reagents		
Sample preservation/ handling		
Procedures	7.3.1 Gasoline Range Organics (GRO): All chromatographic peaks eluting from methyltert-butylether through naphthalene, inclusive. (modify for non-South Carolina clients only)	7.4.2 Two specific gasoline components are used to establish the range, 2-methylpentane and 1,2,4-trimethylbenzene.
Procedures	7.4.1 A gasoline component standard is analyzed instead of a commercial fuel standard. (modify for non-South Carolina clients only).	7.3.3use recently purchased commercially available fuel
QC Method Blank		
QC Accuracy/Precision		

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#### TABLE 3

#### PURGE-AND-TRAP OPERATING PARAMETERS

RECOMMENDED CONDITIONS FOR SUGGESTED TRAP			
Purge gas	Nitrogen or Helium		
Purge gas flow rate (mL/min)	40		
Purge time (min)	11.0 <u>+</u> 0.1		
Purge temperature (°C)	40°C or less		
Desorb temperature (°C)	250°C		

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TABLE 4
GASOLINE STANDARD COMPONENTS AND CONCENTRATIONS

COMPONENT	CONCENTRATION, ug/mL
Methyl-tert-butylether (MTBE)	1000
Benzene	1000
Toluene	1000
Ethylbenzene	1000
m-Xylene	1000
p-Xylene	1000
o-Xylene	1000
1,2,4-Trimethylbenzene	1000
1,3,5-Trimethylbenzene	1000
Naphthalene	1000
TOTAL	10,000

TABLE 5

#### **PQLs**

Analyte	Water	Soil
	ug/L	mg/Kg
GRO	10	2.5

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#### FIGURE 1

#### GC SOIL PREP LOGBOOK

#### KATAHDIN ANALYTICAL SERVICES GC SOIL PREP LOG

Date of Sample Preparation	Analyst Initials	Sample #	Sample Weight (g)	Volume (mL) Methanol  or DI Water  <	Spike ID and Volume (μL)	Surrogate ID and Volume (µL)	Method	Comments
2-17-12	EKC	WG104863-1	10.00	10 mush	_	-	8015m-GRD	
	1	1 -2	9.99	1	1254L AMP3120 @ 2000 usimi	-	1	
		-3	10.62		1	_		
		SF 6324-15	5.00	5 mlx	-	_		XIML LOT SOSS-SLOWL DI
J		SF6437-3	5.00	5 MWH	50-41 457622m56	_		
2-22-12	EKC	WG104971-1	15.00	15 MUDH	NA	25000 majore	MA-VPH	R187899
	j	1 -2	15.02	1	35 ml Am P3099 @ 10,000 ugime	NA	1	
		- 3	15-01		1	+		
		SE0785-1	18.48		NA	2001 AMP3245		
		1 -2	19.49		1			
		-3	16.45					
		SF0804-1	17.52			↓		
		SF0843-1	15.16			0 5000 ughu		
		SF0847-3	8.54					
		1 -4A	8.24	4				State on the state of the state
	1	1 -46	6.48			1		pink

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FIGURE 2

#### **INSTRUMENT RUNLOG PAGE**

Method (	Circle	): MADEP-VPH-98-	1 ME	DEP	4.2.17	(SV	V846	8015	(M)		
Date	Init.	Sample Name	Amt. Purged	SP.	Res		Dil.	Y/N	Method	pH	Comments
2-10-12	MAM		SmL	a	26810	_	1	Y	NOHPTOIS	-	
2-13-12		13	-	_	DEBIO		-	_	- WASTERDIE	_	
1	1	CUSO	SmL	1	N DIO	37	1	Y		NIA	
		WG104470-1842		2		36	1	4		1	
		SF0553-1AA	T	3		39	1	Y		1	
1	1	CV 50	SML	4		40	1	Y	1	1	
2-15-12	EKI.	TB	-	_	1	41	-	N	GR001	-	
1	1	CV 100	5ml	1		42	1	y	ORCOL	-	
		WG104729-1	1	2		43	1			-	
		1 -7		3		44	1			-	
	1	-3		4		45	1			-	
		SED751-14NL	2.5m	5		46	2	N			- 1 - 1 1
		1 -14	Smi *	1		47	1	y		-	10+ nuded 4500 60+ 319-62
		water	5mL	7		48	1	N			ICO ML DI
		AG-mb-l	1	K		49	$\vdash$	7			
		1 2		9		50	$\vdash$	i			2 W 45+47
		3		10		51	+	N		_	
		ч		11		52	Н	Y			and bot bre
		5		12		53	+	1		-	
				13		54	+	H			
		,		14		55	+	H		_	
		8		15		56	+	Н			-
2-16-12		Louk		13		57		H		-	
10-10		AD LON		110		58					4
		AQ LOQ		2		9	+			-	0.9 el #7
		C.V 250		3		0.				_	1-Dul #7
2-17-17	EX.	EV 100 0/1	0217	112	-	.0		4	-		
2-17-12	EXL	TA	- 031	1 1	2FAIC	1.16	_	N	GROOT	-	
1	1	CV 100	SML	)	7	62	1	7	1	-	
Std. Na	-	0		011						1	
VPH Cal	mix	Conc. 50 ug/ml		Std. 0			Std. (				CC = 5uL Std. 1 LCS = 5uL Std. 3
VPH Surr	. mix	100 ug/ml	GCV 7	185	8		GCV				Samples = 5uL Std. 2
GRO Cal	mix		GCV 3	385	9		GCV GCV			GRO	CC = 2uL Std. 4
GRO Sun	r. Mix	50 ug/mL	GCV 25	352			GCV				LCS = 2uL Std. 4
GRO LCS		250 ug/mL Total	GCV 29	51.0			GCV			GRO	Samples = 2uL Std. 5
morner	ne	50 uglml	GCM 28	46		_					

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#### FIGURE 3

#### PRIMARY REVIEW CHECKLIST

#### PRIMARY REVIEW CHECKLIST

nt:		Primary	Secondary
thod:		Date:	Date:
G No:	Level:	Initials:	Initials:
S No:			Approved :
DODQSM  List all curv	( REPORT ND's to		QUAPP □ LAB □ <u>LOD</u> □)
	ich QC limits were us	sed for ( Surr., LCS's MS/M	SD's.)
Correct Wo	rk Order Number or S	DG name (all forms).	
Correct proj	ect name and spelling	(all forms). (Truncated $\square$	).
Correct file	numbers (all forms).		-
Analysis Da	ate Correct.		-
Extraction N	Method & Analysis M	ethod Correct.	
Product list	compared to ROAs (c	compounds & PQLs).	
Chromatogr	ram reviewed for unlal	beled peaks (check product	list).
Flagging of	all ROAs correct (F	lorida Flagging 🗆 ).	1
All tunes in	cluded (level IV).		1 <del>7 </del>
All log bool	c pages included (Soil	weights, TCLP & SPLP).	Y
Verify DOI	O QSM criteria and/or	Project specific requirement	nts
Narrate any	y method deviations.	(Blanks, LCS's etc.)	
Sign & Date	Manual integration	( Narrate as needed ).	
Camala I D	's Truncated (NARI	DATED VEC D	ease list KAS # below :

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#### KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-502 **Revision History Cover Page** Page 1

TITLE: PREPAR ANALYS	RATION OF AQUEOUS SAMPLES FOR EXTR	ACTABLE	SEMIVOLATILE
Prepared By:	Micheal Thomas	Date:	07-24-00
Approved By:			
Department Manage	er: 1 two	Date:	6-23-06
Operations Manage	r. Actorah Madeau	Date:	6.23.00
QA Officer:	Liseie Dimond	Date: <u>_</u>	6-23-06

Revision History:

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
03	Changes to sect. S.S : Figures 3:4 to refield corrent spike solutions and concentrations Replaced cover page.  Original cover page filed with SOP CASOZ-02	LAO	04/06	04/06
04	Added definitions, added waste information added LCSD, added SIM LCS/D, ms/D, updated Table 1, added use of narrow range pH paper. Minor changes throughout to reflect current	LAD	०९(०२	09/07
05	Minor changes throughout to reflect current Removed ms/mso 14 day requirements. Changed CLLE extraction time to 18 > 24 hours. Added information on determining initial Sample volume. Added extracted Sample Hisposal. Removed all references to method 625.	UAD	09108	09/08
06	Added to check PH ofter BIN CLLE extraction to ensure pH >11. If not add more Madand continue extracting. Added information for initial Volume Jetermination. Added Reference to CA-108. updated losbook example. Added if extract goes dry-veextract	LAYD	10/09	10/09
07	Sect. 5 - Removed backing and rinsing NaSO4. Added 1.4-Dioxane to SIM surrogete Mix, Sect. 7 added acid to BIN SIM., removed to let separate for 10 minutes, minor edits throughout.	LAD	03/12	03/12

# KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-502 Revision History Cover Page (cont.) Page 2

TITLE:	PREPARATION OF AQUEOUS SAMPLES FOR EXTRACTABLE SEMIVOLATILE
	ANALYSIS

Revision History (cont.):

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
08	Removed Sect. 7.1.9, determining the sample initial volume. Sect. 7.1.4 has this information. Figures 1 and 2 updated.	UND	05/13	05/13
	-			

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TITLE:	PREPARATION OF AQUEOUS SAMPLES FOR E	EXTRACTABLE SEMIVOLATILE				
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# TITLE: PREPARATION OF AQUEOUS SAMPLES FOR EXTRACTABLE SEMIVOLATILE ANALYSIS

#### 1.0 SCOPE AND APPLICATION

The purpose of this SOP is to describe procedures utilized by Katahdin Analytical personnel in the preparation of all non-CLP aqueous samples for analysis of extractable semivolatile organic compounds.

The goal of this procedure is to ensure uniformity involving the preparation of samples for subsequent SVOA analysis by GC/MS. This SOP is applicable to EPA Methods 3510 (modified separatory funnel extraction) and 3520 (continuous liquid-liquid extraction), current revisions.

#### 1.1 Definitions

METHOD BLANK (laboratory reagent blank): An artificial sample designed to determine if method analytes or other interferences are present in the laboratory environment, the reagents, or the apparatus. For aqueous samples, reagent water is used as a blank matrix; for soil samples, baked organic-free sand is used as a blank matrix. The blank is taken through the appropriate steps of the process.

LABORATORY CONTROL SAMPLE (LCS): A blank that has been spiked with the analyte(s) from an independent source, and is analyzed exactly like a sample. Its purpose is to determine whether the methodology is in control, and whether the laboratory is capable of making accurate and precise measurements. The matrix used should be phase matched with the samples and well characterized.

MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD): Predetermined quantities of stock solutions of certain analytes are added to a sample matrix prior to sample extraction and analysis. Samples are split into duplicates, spiked and analyzed. Percent recoveries are calculated for each of the analytes detected. The relative percent difference between the samples is calculated and used to assess analytical precision.

SURROGATES: Organic compounds which are similar to analytes of interest in chemical composition, extraction and chromatography, but which are not normally found in environmental samples. These compounds are spiked into all blanks, standards, samples and spiked samples prior to analysis. Percent recoveries are calculated for each surrogate.

#### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of analysts experienced in the extraction of samples for semivolatile analysis. Each analyst must demonstrate and document their ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, current revision, "Personnel Training & Documentation of Capability".

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## TITLE: PREPARATION OF AQUEOUS SAMPLES FOR EXTRACTABLE SEMIVOLATILE ANALYSIS

It is the responsibility of all Katahdin technical personnel involved in the extraction of samples for semivolatile analysis to read and understand this SOP, to adhere to the procedures outlined, and to properly document their data in the appropriate lab notebook. Any deviations from the test or irregularities with the samples should also be recorded in the lab notebook and reported to the Department Manager or designated qualified data reviewer responsible for this data.

It is the responsibility of the Department Manager to oversee that members of their department follow this SOP, to ensure that their work is properly documented and to initiate periodic review of the associated logbooks.

#### 1.3 Safety

Users of this procedure must be cognizant of inherent laboratory hazards, proper disposal procedures for contaminated materials and appropriate segregation of hazardous wastes. The toxicity or carcinogenicity of each reagent used in this method has not been precisely defined; however, each chemical should be treated as a potential health hazard. A reference file of material safety data sheets is available to all personnel involved in the chemical analysis. Everyone involved with the procedure must be familiar with the MSDS's for all the materials used in this procedure.

Each qualified analyst or technician must be familiar with Katahdin Analytical Environmental Health and Safety Manual including the Katahdin Hazardous Waste Management Plan and must follow appropriate procedures. These include the use of appropriate personal protective equipment (PPE) such as safety glasses, gloves and lab coats when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location of all safety equipment. Each analyst shall receive a safety orientation from their Department Manager, or designee, appropriate for the job functions they will perform.

#### 1.4 Pollution Prevention/Waste Disposal

Whenever possible, laboratory personnel should use pollution prevention techniques to address their waste generation. Refer to the current revision of the Katahdin Hazardous Waste Management Plan for further details on pollution prevention techniques.

Wastes generated during the preparation of samples must be disposed of in accordance with the Katahdin Hazardous Waste Management Plan and Safety Manual and SOP SD-903, "Sample Disposal," current revision. Expired standards are lab packed, placed in the Katahdin hazardous waste storage area, and disposed of in accordance with this SOP.

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Any methylene chloride solvent waste generated during the rinsing of glassware, disassembly of CLLEs after extraction, etc. should be disposed of in the "D" waste stream satellite accumulation area nearest the point of generation. Acetone and methanol are considered flammable waste, and should be disposed of in the "O" waste stream satellite accumulation area nearest the point of generation. Post-extraction aqueous samples are considered either N-Hi or N-Low waste and should be disposed of in the corresponding satellite waste accumulation area nearest the point of generation. Sodium sulfate used for sample drying should be disposed of in the soil with organics "I" waste stream satellite accumulation area nearest the point of generation. Please refer to the current revision of SOP CA-107 for the location of satellite waste accumulation areas.

#### 2.0 SUMMARY OF METHOD

For aqueous samples extracted by CLLE, a one liter aliquot of sample is adjusted to pH  $\leq$  2 and extracted with methylene chloride using a continuous liquid-liquid extractor. The pH is then adjusted to pH  $\geq$  11 and the sample is extracted again with methylene chloride. A modified separatory funnel extraction may also be used. If this procedure is used, the sample aliquot is first adjusted to pH  $\geq$  11 and then to pH  $\leq$  2. The methylene chloride extract is dried and concentrated to a volume of 1.0 mL.

#### 3.0 INTERFERENCES

Solvents, reagents, glassware, and other sample preparation apparatus may yield interferences to GC/MS analysis due to the presence of contaminants. These contaminants can lead to discrete artifacts or elevated baselines in the total ion current profiles (TICPs). Routinely, all of these materials must be demonstrated to be free from interferences under the conditions of the analysis by running reagent blanks. Interferences caused by phthalate esters can pose a major problem in semivolatile analysis. Common flexible plastics contain varying amounts of phthalates that are easily extracted during laboratory operations, so cross-contamination of glassware frequently occurs when plastics are handled. Interferences from phthalates can best be minimized by avoiding the use of such plastics in the laboratory. At no time may gloves that have not been tested for phthalates or gloves known to contain phthalates be used or stored in the organic extraction lab. Additionally, whenever possible plastic items in this lab must be replaced with metal or teflon or other non-phthalate plastic substitute.

Special care should be taken to ensure that clean glassware and apparatus are used and pre-rinsed with the appropriate solvent prior to use. Solvents should be analyzed prior to use to demonstrate that each lot is free of contaminants that may interfere with the analysis.

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Interferences coextracted from the samples will vary considerably from source to source. If analysis of an extracted sample is prevented due to inteferences, further cleanup of the sample extract may be needed to minimize interferences.

#### 4.0 APPARATUS AND MATERIALS

Brand names and catalog numbers are included for illustration purposes only.

- 4.1 Continuous liquid-liquid extractors including body, 500 mL round bottom flask and Alhin condensers and equipped with Teflon or glass connecting joints requiring no lubrication (Hershberg-Wolf Extractor, Ace Glass Company, Vineland, NJ, P/N 6841-10 or equivalent).
- 4.2 Glass powder funnels.
- 4.3 Fluted filter paper, 18.5cm diameter.
- 4.4 Concentrator tube Kuderna-Danish, 10 mL, graduated (Kontes K-570050-1025 or equivalent). Calibration must be checked at the volumes employed in the test.
- 4.5 Evaporation flask Kuderna-Danish, 500 mL (Kontes K-570001-0500 or equivalent). Attach to concentrator tube with neck clips.
- 4.6 Snyder column Kuderna-Danish, three- or four-ball macro (Kontes K-503000-0121 or equivalent).
- 4.7 Syringe gas tight, 1.0 mL, solvent rinsed between each use.
- 4.8 Vials Glass, 1.8 mL capacity, with polytetrafluoroethylene (PTFE)-lined screw top and 12 mL with Teflon-lined caps.
- 4.9 2 L separatory funnel, equipped with Teflon stopper and stopcock; Nalgene Teflon FEP separatory funnels may also be used.
- 4.10 Organic Free Boiling Chips approximately 10/40 mesh, Teflon or silicon carbide (or equivalent). Cleaned by Soxhlet for 18 hours.
- 4.11 Water bath heated, with concentric ring cover, capable of temperature control (± 20°C). The bath should be used in a hood.
- 4.12 Nitrogen evaporation apparatus.
- 4.13 Wide range pH test strips, pH 0-14, Whatman CF Type.

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- 4.14 Glass rods for stirring samples.
- 4.15 Amber bottles or other appropriate containers for collection of extracts from separatory funnel extraction.
- 4.16 5 3/4" Pasteur pipets.
- 4.17 Narrow range pH test strips, pH 0 to 2.5 pH, EMD ColorpHast or equivalent.
- 4.18 Narrow range pH test strips, pH 11 to 13 pH, EMD ColorpHast or equivalent.

#### 5.0 REAGENTS

All reagent and solvent lots must be checked for possible contamination. Refer to the current version of Katahdin SOP CA-105, Reagent and Solvent Handling, for further details. The extraction staff is responsible for submitting samples to the GC or GC/MS sections for appropriate analysis. All information concerning preparation of the reagent/solvent lot sample will be recorded in the Organic Extraction Log (Figure 1) and acceptance or rejection of these lots must be recorded in the solvent/reagent lot check logbook (Figure 2). All reagents and solvents must be free (<PQL) of any target compounds.

- 5.1 <u>Laboratory Reagent Grade Water</u> defined as water in which an interferent is not observed at or above the PQL of each parameter of interest. Deionized water filtered through activated charcoal.
- 5.2 <u>Sodium sulfate</u> (ACS reagent grade) granular, anhydrous, certified by the manufacturer/vendor as purified.
- 5.3 Sulfuric acid solution (1:1  $H_2SO_4$ :  $H_2O$ ) Prepared in an icebath by slowly adding a volume of concentrated  $H_2SO_4$  to an equivalent volume of reagent water and swirl gently to mix. Caution should be taken when adding the acid to the water as the reaction is highly exothermic.
- 5.4 <u>Acetone, methanol, methylene chloride</u> pesticide residue analysis grade or equivalent, evaluated prior to use by concentration of 200 mLs to 1.0 mL followed by GC and/or GC/MS analysis.
- 5.5 <u>Standard Preparation</u> For all standard preparations, see current revision of the following Katahdin Analytical SOPs:
  - "Standards Preparation, Documentation and Traceability", (CA-106, current revision)

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- "Balance Calibration," (CA-102, current revision)
- 5.5.1. Base/Neutral and Acid (SVOA) Surrogate Spiking Solution Surrogate standards are added to all samples and calibration solutions. Prepare a surrogate standard spiking solution that contains the following compounds at the indicated concentrations in acetone.

Compound	Conc.
phenol- <sub>d6</sub>	100 ug/mL
2,4,6-tribromophenol	100 ug/mL
2-fluorophenol	100 ug/mL
nitrobenzene- <sub>d5</sub>	50 ug/mL
p-terphenyl- <sub>d14</sub>	50 ug/mL
2-fluorobiphenyl	50 ug/mL

Store the spiking solution at -10°C to -20°C in Teflon-sealed containers. These solutions must be replaced after six months, or sooner if comparison with quality control check samples indicates a problem. If reanalysis of a method blank still indicates surrogates out of criteria, a new surrogate solution must be used.

5.5.2 SIM Surrogate Spiking Solution- Surrogate Standards are added to all samples and calibration solutions. Prepare a surrogate solution that contains the following compounds at a concentration of 2 ug/mL in acetone.

Compound	Conc. ug/mL
Fluorene-d10	2.0 ug/mL
2-Methylnaphthalene-d10	2.0 ug/mL
Pyrene-d10.	2.0 ug/mL
2,4-Dibromophenol	4.0 ug/mL
1,4-Dioxane-d8	20 ug/mL

Store the spiking solution at -10°C to -20°C in Teflon-sealed containers. These solutions must be replaced after six months, or sooner if comparison with quality control check samples indicates a problem. If reanalysis of a method blank still indicates surrogates out of criteria, a new surrogate solution must be used.

5.5.3 SVOA Matrix Spike/Lab Control Samples Spiking Solution - the matrix spike/LCS solution consists of the compounds listed in Figure 3. Prepare a spiking solution that contains each of the base/neutral compounds listed in Figure 3 at 50 ug/mL in methanol and the acid compounds at 100 ug/mL in methanol. Matrix spike/LCS standards are stored in the freezer (-10°C to -20°C) located in the storage area.

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- 5.5.4 Base/Neutral/Acid (SIM) Matrix Spike/ Lab Control Sample Spike Solution for SIM-SVOA. Prepare a spiking solution in methanol that contains the compounds listed in Figure 3 at a concentration of 2 ug/mL for base/neutrals and 4.0 ug/mL for acids. Take out 1.0 mL of Base/Neutral and Acid Matrix Spike/Lab Control Spiking Solution for SVOA and dilute it to 25.0 mL of methanol. Store the solution Spiking at -10°C to -20°C in Teflon-sealed containers. These solutions must be replaced after six months or sooner if comparison with quality control check samples indicates a problem.
- 5.5.5 Base/Neutral and Acid (SVOA) Appendix IX Lab Control Sample / Matrix Spike Spiking Solution Prepare a spiking solution in methanol that contains the compounds listed in Figure 4 at concentrations of 100 ug/ml. Store the spiking solution at -10°C to -20°C in Teflon-sealed containers. These solutions must be replaced after six months, or sooner if comparison with quality control check samples indicates a problem.

#### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

Continuous liquid-liquid (Method 3520) and/or separatory funnel (Method 3510) extractions for semivolatiles must be started within seven days of date of sample collection, although the analyst should be aware that actual holding times employed may be project/program specific. If sampling date is unknown, the hold time is counted from one day prior to date received.

#### 7.0 PROCEDURES

The following information must be recorded in the extraction logbook.

- Extraction method
- Surrogate and spike IDs
- Lot numbers of all solvents, acids and bases, sodium sulfate, filter paper
- Nitrogen evaporation water bath temperature
- Sample pH if applicable
- Extraction and Concentration dates
- Extraction and Concentration analyst
- Sample ID or QC sample ID
- Initial and final volumes or weight
- Surrogate and spike amounts
- Any sample cleanup preformed
- Final extract tray location
- Any comments regarding the sample extraction (ie. Emulsion)

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- Prep batch start time and end time
- CLLE start time and end time
- Lot number of the vials the concentrated extracts are stored in.

Follow the proper procedures for maintaining Internal Chain of Custodies for samples when removing and replacing samples in storage locations. This procedure is described in KAS SOP SD-902, "Sample Receipt and Internal Control", current revision.

- 7.1 CONTINUOUS LIQUID-LIQUID EXTRACTION (Method 3520)
  - 7.1.1 Set up the CLLE apparatus. All glassware should be pre-rinsed three times with methylene chloride in order to eliminate any contamination factors.
  - 7.1.2 Add approximately 500 600 mL of methylene chloride to the CLLE body. Label each flask with the following: sample number (or QC identification number), analyte (SVOA), extraction method (CLLE), and extraction date.
  - 7.1.3 A method blank and a laboratory control sample (LCS) must be prepared for each daily extraction batch of twenty samples or fewer (if a work order consists of more than twenty samples, a new batch must be started on a separate page with its own method blank and LCS). To prepare method blank and LCS, add 1 L reagent water to a CLLE body. Be sure that no water leaks into the round bottom flask. If combined SIM-SVOA analysis is requested, a separate LCS must be prepared for each analysis. This blank and LCS are carried through the entire extraction and analytical procedure.
  - 7.1.4 Measure the initial volume by comparing the meniscus of the sample with the reference bottle of the same bottle type. Please refer to SOP CA-108, "Basic Laboratory Technique", for the reference bottle verification procedure. Record the volume and any notable characteristics (e.g. color, presence of sediment, or odor) in the extraction logbook.
  - 7.1.5 Transfer the sample to a CLLE body slowly, being sure that no water leaks into the round bottom flask.
  - 7.1.6 If the batch requires a MS/MSD, transfer two 1 L portions of the sample selected/designated for MS/MSD to CLLE bodies for preparation of a matrix spike/matrix spike duplicate if required. If combined SIM-SVOA analysis is requested, a separate MS/MSD must be prepared for each analysis. If extra MS/MSD aliquots of sample are unavailable a laboratory control sample duplicate (LCSD) may be substituted.

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- 7.1.7 Check the pH of each sample with wide range pH paper by removing a couple of sample drops with a clean disposable pipet or on the tip of a stirring rod. Adjust the pH of the samples (including method blank, LCS/LCSD, and MS/MSD) to  $\leq$  pH 2 with 1:1 H<sub>2</sub>SO<sub>4</sub> after addition of surrogates and spikes and prior to attaching Allihn condensers (Step 7.1.11). Stir with a glass stirring rod and check pH by tapping the glassrod onto wide range pH paper. The pH must be  $\leq$  2. If the pH test strip does not clearly indicate the pH is less than 2, narrow range pH paper must be used.
- 7.1.8 For each sample, rinse the original sample container with approximately 30 mL of methylene chloride. Add this rinse to the CLLE body.
- 7.1.9 To all samples, method blank, LCS/LCSD, and MS/MSD add 1.0 mL base/neutral and acid (SVOA) surrogate spiking solution using a 1.0 mL or 2.5 mL gas tight syringe. Record surrogate spike volume and identification code in extraction logbook. Thoroughly rinse syringe with acetone before and after each use.
  - 7.1.9.1 If the request is for SVOA, use the SVOA Surrogate Solution (sect. 5.5.1).
  - 7.1.9.2 If the request is for SIM, use the SIM surrogate solution (sect. 5.5.2).
  - 7.1.9.3 If the request is for SIM-SVOA, use the SIM surrogate solution as well as SVOA surrogate solution. NOTE: Separate LCS/LCSD and/or MS/MSD are needed for each analysis and should be spiked with the appropriate surrogate.
- 7.1.10 To LCS/LCSD and MS/MSD add 1.0 mL base/neutral and acid (SVOA) matrix spike/LCS spiking solution using a 1.0 mL or 2.5 mL gas tight syringe. Record matrix spike/LCS spiking solution volume and identification code in extraction logbook. Thoroughly rinse syringe with methanol before and after each use.
  - 7.1.10.1 If the request is for SVOA add 1.0 mL of SVOA Matrix Spike/Lab Control Samples Spiking Solution (sect 5.5.3).
  - 7.1.10.2 If the request is for SIM add 1.0 mL of SVOA Matrix Spike/Lab Control Samples Spiking Solution (sect 5.5.3) and add 1.0 mL of Base/Neutral/Acid (SIM) Matrix Spike/ Lab Control Sample Spike Solution (sect 5.5.4).
  - 7.1.10.3 If the request is for SVOA Appendix IX, use the SVOA Appendix IX Spiking solution as well as the SVOA spiking solution -add 1.0 mL of

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SVOA Matrix Spike/Lab Control Samples Spiking Solution (sect 5.5.3) and add 1.0 mL of Base/Neutral and Acid (SVOA) Appendix IX Lab Control Sample / Matrix Spike Spiking Solution (sect 5.5.5).

- 7.1.11 Attach cooling water Allihn condensers, after first rinsing each 45/50 joint with methylene chloride. Turn on the heating mantles and allow the samples to extract for 18 to 24 hours. Turn off the mantles and let samples cool.
- 7.1.12 Detach condensers and verify that the pH is still  $\leq 2$  in the same manner mentioned in 7.1.6. If the pH has changed, more acid should be added to make the pH  $\leq 2$  and the sample extracted for several more hours.
- 7.1.13 Upon completion of acid extraction, allow the sample to cool. Detach condensers and add enough 10N NaOH to adjust the pH to ≥ 11 with stirring. Use glass stirring rods to stir and check the pH of each sample in the same manner mentioned in 7.1.6.
- 7.1.14 Re-attach Allihn condensers, turn on heating mantles, and allow samples to extract for 18 to 24 hours. Turn off mantles and allow samples to cool.
- 7.1.15 Detach condensers and verify that the pH is still  $\geq$  11 in the same manner mentioned in 7.1.6. If the pH has changed, more NaOH should be added to make the pH  $\geq$  11 and the sample extracted for several more hours.
- 7.1.16 Once samples are cool to the touch, the CLLE apparatus can be disassembled. The round bottom flask is removed, covered foil and placed in the interim extract refrigerator. The remaining sample in the CLLE body is poured in the "N-Hi" satellite.
- 7.1.17 Proceed to Step 7.3 for sample extract concentration procedures.
- 7.2 SEPARATORY FUNNEL EXTRACTION (Modified Method 3510)

If an emulsion prevents acceptable recovery or client history indicates samples may demonstrate matrix interference, then samples should be extracted by continuous liquid-liquid extraction (CLLE).

- 7.2.1 Rinse <u>all</u> glassware, including teflon separatory funnels, three times with methylene chloride prior to use.
- 7.2.2 Label 2 L separatory funnels and amber collection bottles clearly. Each label should include: sample number (or QC indicator number), analyte (SVOA), matrix (Aq), extraction date.

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- 7.2.3 A method blank and a laboratory control sample (LCS) must be prepared for every 20 samples or with each extraction batch, whichever is more frequent. To prepare method blank and LCS, add 1 L reagent water to a separatory funnel. If combined SIM-SVOA analysis is requested, a separate LCS must be prepared for each analysis. This blank and LCS are carried through the entire extraction and analytical procedure.
- 7.2.4 Measure the initial volume by comparing the meniscus of the sample with the reference bottle of the same bottle type. Please refer to SOP CA-108, "Basic Laboratory Technique", for the reference bottle verification procedure. Record the volume and any notable characteristics (e.g. color, presence of sediment, or odor) in the extraction logbook.
- 7.2.5 If the batch requires a MS/MSD, transfer two 1 L portions of the sample selected/designated for MS/MSD to separatory funnels for preparation of a matrix spike/matrix spike duplicate if required. If combined SIM-SVOA analysis is requested, a separate MS/MSD must be prepared for each analysis. If extra MS/MSD aliquots of sample are unavailable, a laboratory control sample duplicate (LCSD) may be substituted.
- 7.2.6 To all samples, method blank, LCS/LCSD, and MS/MSD add 1.0 mL base/neutral and acid (SVOA) surrogate spiking solution using a 1.0 mL or 2.5 mL gas tight syringe. Record surrogate spike volume and identification code in extraction logbook. Thoroughly rinse syringe with acetone before and after each use.
  - 7.2.6.1 If the request is for SVOA, use the SVOA Surrogate Solution.
  - 7.2.6.2 If the request is for SIM, use the SIM Surrogate Solution.
  - 7.2.6.3 If the request is for SIM-SVOA, use the SIM surrogate solution as well as SVOA surrogate solution. NOTE: Separate LCS/LCSD and/or MS/MSD are needed for each analysis and should be spiked with the appropriate surrogate.
- 7.2.7 To LCS/LCSD and MS/MSD add 1.0 mL base/neutral and acid (SVOA) matrix spike/LCS spiking solution using a 1.0 mL or 2.5 mL gas tight syringe. Record matrix spike/LCS spiking solution volume and identification code in the extraction logbook. Thoroughly rinse syringe with methanol before and after each use.
  - 7.2.7.1 If the request is for SVOA, use the SVOA Spiking Solution.
  - 7.2.7.2 If the request is for SIM, use the SIM Spiking solution.

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- 7.2.7.3 If the request is for SVOA Appendix IX, use the SVOA Appendix IX Spiking solution as well as the SVOA spiking solution
- 7.2.8 For each sample, rinse the original sample container with 60 mL of methylene chloride. Add this rinse to the separatory funnel.
- 7.2.9 Adjust the pH of the samples (including method blank, LCS/LCSD, and MS/MSD) to pH ≥ 11 with 10N NaOH after addition of surrogates and spikes. Stir with a glass stirring rod and check pH by tapping the glass stirring rod onto wide range pH paper. The pH must be ≥ 11. If the pH test strip does not clearly indicate the pH is greater than 11, narrow range pH paper must be used.
- 7.2.10 Add 60 mL of methylene chloride directly to the method blank and LCS/LCSD separatory funnels.
- 7.2.11 Extract the samples by shaking the funnel for two minutes, venting often, but gently, in a hood to release pressure. A mechanical shaker may be used, where samples are shaken for 3 minutes. Following each shake, allow phases to separate. Drain the methylene chloride layer into an amber collection bottle.
- 7.2.12 If an emulsion forms, mechanical techniques must be employed to achieve maximum separation. Such means include swirling, centrifugation, and draining through a small separatory funnel. In certain instances, transferring the entire sample into a continuous liquid-liquid extractor (CLLE) may be the only alternative. If any such techniques are used, they must be noted in the extractions logbook, and the batch transferred to a CLLE batch with its own batch ID.
- 7.2.13 Add a second 60 mL aliquot of methylene chloride to the separatory funnel and extract for the second time (see 7.2.12 7.2.13). Collect the methylene chloride layer in the same amber collection bottle.
- 7.2.14 Repeat the extraction for a third time as described in 7.2.14.
- 7.2.15 Following the third shake, using a glass stirring rod, check the pH to ensure that it has remained at  $\geq$  11. If the pH has changed back to neutral range, it must be readjusted to  $\geq$  11 and the sample must be extracted at least one more time, adding the methylene chloride to the same amber bottle, that was previously used. If the pH has remained at a value  $\geq$  11, the pH is then adjusted to  $\leq$  2 with 1:1 H<sub>2</sub>SO<sub>4</sub>. Add enough 1:1 H<sub>2</sub>SO<sub>4</sub> to adjust the pH to  $\leq$  2 with stirring. Use glass stirring rods to stir.

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- 7.2.16 Add 60 mL methylene chloride and extract the samples three times in the same manner described in 7.2.11 7.2.13. Collect the methylene chloride layer in the same amber collection bottle used to collect the acid fraction.
- 7.2.17 Sample waste should be poured into the "n-lo" satellite.
- 7.2.18 Proceed to Section 7.3 for extract concentration procedures.

#### 7.3 CONCENTRATING THE EXTRACTS

- 7.3.1 For Methods 3510 and 3520, the combined fractions are concentrated to a final volume of 1.0 mL.
- 7.3.2 Rinse the K-D glassware (flask, concentration tube, and snyder column, including the ground glass joints on the flask and columns) three times with methylene chloride. Add two boiling chips to the K-D prior to final rinse. Also rinse the assembled funnels, filter paper, and granular sodium sulfate used for drying the extracts.
- 7.3.3 Transfer the methylene chloride extract to a K-D concentrator setup through a short stem funnel filled with 1-2 inches of sodium sulfate in filter paper. This is the drying step, which is required to remove residual water from the extracts. Any large water layers must be removed by other means, prior to pouring through the sodium sulfate. After pouring all of the extract volume through the sodium sulfate, rinse the extract flask three times with  $\sim 2-3$  mls of methylene chloride. Add the rinsings through the sodium sulfate to complete a quantitative transfer. Rinse the sodium sulfate with  $\sim 15$  mls of methylene chloride and allow to drain
- 7.3.4 Transfer the label from the collection bottle or round bottom flask (for CLLE) to a K-D. Remove the funnel and attach a 3- or 4-ball macro Snyder column. Pre-wet the Snyder column with 1 mL of methylene chloride.
- 7.3.5 Place the K-D in a hot water bath (75-85°C). Gently swirl K-D in the water until boiling begins. At the proper distillation rate, the Snyder balls should chatter but the chambers should not flood with condensed solvent. The K-D should be kept in a vertical orientation while on the bath. When the apparent volume in the concentrator tube reaches  $\approx 6$  mL, remove the K-D from the water bath. Allow the K-D to cool for 10 minutes. Rinse the Snyder column lower joint with  $\approx 1$  mL of methylene chloride. Remove the Snyder column. Wipe off any water from the neck above the lower joint of the flask. Separate the K-D flask from the concentrator tube, rinsing the ground glass joint with  $\approx 1$  mL methylene chloride.

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# TITLE: PREPARATION OF AQUEOUS SAMPLES FOR EXTRACTABLE SEMIVOLATILE ANALYSIS

- 7.3.6 Reduce the methylene chloride extract in the concentrator tube to approximately 1 mL using the nitrogen blow-down apparatus. The bath temperature must be no higher than the boiling point of the solvent (39°C for methylene chloride). Turn the gas to 3 psi. Be careful not to splash the extract out of the tube. During concentration on the N-evap, the internal wall of the concentrator tube and the N-evap sparging pipet must be rinsed down at least once or twice with ≈1 mL of methylene chloride. The solvent level in the concentrator tube must be positioned below the level of the water bath as much as possible to prevent water from condensing into the sample extract. As the extract volume is reduced, lower the N₂ sparging pipet closer to the surface of the extract to expedite the concentration. Note any problems or extract losses, if they occur, in the extractions logbook.
- 7.3.7 Reduce each extract to slightly less than 1 mL and then, using a 5 ¾" pasteur pipet, transfer the final extract and label to a 1.8 mL vial with PTFE-lined cap.
- 7.3.8 If at any time during the concentration process the concentrator tube goes dry, reextraction must occur immediately.
- 7.3.9 Transfer all of the extract to a 1.8 mL screw cap vial. Using methylene chloride, adjust the final volume of each extract to 1 mL by comparison to an appropriate reference vial.

Store in refrigerator until GC/MS analysis.

#### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

A method blank must be extracted for each and every item listed below:

- Each sample matrix (soil, water)
- Each day of extraction (24 hours midnight midnight)
- Each extraction method or level
- Every 20 samples extracted in a 24-hour period

A laboratory control sample (LCS) is required for <u>each</u> and <u>every</u> item listed below:

- Each sample matrix
- Each extraction method or level
- Every extraction batch of twenty or fewer samples

Refer to the current revision of the applicable Katahdin SOP for analysis of semivolatiles for quality control acceptance criteria.

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# TITLE: PREPARATION OF AQUEOUS SAMPLES FOR EXTRACTABLE SEMIVOLATILE ANALYSIS

#### 9.0 METHOD PERFORMANCE

Refer to the applicable analytical SOP.

#### 10.0 APPLICABLE DOCUMENTS/REFERENCES

Test Methods for Evaluating Solid Waste, Physical/Chemical Methods, USEPA SW846, 3<sup>rd</sup> Edition, Final Updates I, II, IIA, IIB, III, IIIA, IIIB and IV, February 2007, Methods 3510 and 3520, current revisions.

#### LIST OF TABLES AND FIGURES

Table 1	Summary of Method Modifications
Figure 1	Example of Semivolatiles Logbook Page
Figure 2	Example of Solvent/Reagent Lot Check Logbook Page
Figure 3	LCS/Matrix Spike Component List
Figure 4	Appendix Ix LCS/Matrix Spike Component List

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# TITLE: PREPARATION OF AQUEOUS SAMPLES FOR EXTRACTABLE SEMIVOLATILE ANALYSIS

TABLE 1
SUMMARY OF METHOD MODIFICATIONS (METHOD 3510, current revision)

TOPIC	KATAHDIN SOP CA-502-08	METHOD 3510, current revision
Apparatus/Materials Reagents	1) 250 mL amber bottle or flask     2) 1.0 mL syringe     3) short stem funnels	250 mL Erlenmeyer flask     5.0 mL syringe     drying columns
Sample preservation/ handling		
Procedures	<ol> <li>extract collection in amber bottle or Erlenmeyer flask</li> <li>Add surrogate/spike to sample in CLLE</li> <li>Extract for 3 minutes on mechanical shaker</li> <li>extract three times at pH ≥ 11, then extract three times at pH ≤ 2.</li> <li>extract dried using Na<sub>2</sub>SO<sub>4</sub> in short stem funnels</li> <li>Rinse the extract flask three times with ~ 2 - 3 mLs of methylene chloride then rinse the sodium sulfate with ~ 15 mLs of methylene chloride to complete a quantitative transfer</li> <li>water bath temp 75-85 deg C</li> <li>no apparatus height specification for concentration on water bath</li> <li>sample removed from water bath when volume reaches ~6 mL</li> <li>N bath temp no higher than 39 deg C</li> </ol>	<ol> <li>extract collection in Erlenmeyer flask</li> <li>Add surrogate/spike directly to sample bottle</li> <li>Extract by shaking vigorously for 1 - 2 minutes with periodic venting</li> <li>extract three times at pH ≤ 2, then extract three times at pH ≥ 11.</li> <li>extract dried using Na<sub>2</sub>SO<sub>4</sub> in drying columns</li> <li>Rinse the Erlenmeyer flask, which contained the solvent extract, with 20 - 30 mL of methylene chloride to complete the quantitative transfer</li> <li>water bath temp 15-20 deg C above solvent boiling temp</li> <li>partially immerse concentrator tube in water and lower apparatus to complete concentration in 10-20 min</li> <li>sample removed from water bath when volume reaches 1 mL</li> <li>N bath temp 35 deg C</li> </ol>
QC - Spikes	Acid surrogate and spike components at 100 ug/mL; base/neutral surrogate and spike components at 50 ug/mL	Acid surrogate and spike components at 200 ug/mL; base/neutral surrogate and spike components at 100 ug/mL
QC - LCS	Acid surrogate and spike components at 100 ug/mL; base/neutral surrogate and spike components at 50 ug/mL	Acid surrogate and spike components at 200 ug/mL; base/neutral surrogate and spike components at 100 ug/mL

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# TITLE: PREPARATION OF AQUEOUS SAMPLES FOR EXTRACTABLE SEMIVOLATILE ANALYSIS

#### TABLE 1, continued

#### SUMMARY OF METHOD MODIFICATIONS (METHOD 3520, current revision)

TOPIC	KATAHDIN SOP CA-502-08	METHOD 3520, current revision
Apparatus/Materials	short stem funnels	drying columns
Reagents		
Sample preservation/ handling		
Procedures	<ol> <li>Add surrogate/spike to sample in CLLE</li> <li>Add approximately 500 - 600 mL of methylene chloride to the CLLE body</li> <li>CLLE for 22 ± 2 hours</li> <li>Extract dried using Na<sub>2</sub>SO<sub>4</sub> in short stem funnels</li> <li>Rinse the extract flask three times with ~ 2 - 3 mLs of methylene chloride then rinse the sodium sulfate with ~ 15 mLs of methylene chloride to complete a quantitative transfer</li> <li>water bath temp 75-85 deg C</li> <li>no apparatus height specification for concentration on water bath</li> <li>sample removed from water bath when volume reaches ~6 mL</li> <li>N bath temp no higher than 39 deg C</li> </ol>	<ol> <li>Add surrogate/spike directly to sample bottle</li> <li>Add 300 - 500 mL of methylene chloride to the distilling flask of the extractor</li> <li>CLLE for 18 - 24 hours</li> <li>Extract dried using Na<sub>2</sub>SO<sub>4</sub> in drying columns</li> <li>Rinse the Erlenmeyer flask, which contained the solvent extract, with 20 - 30 mL of methylene chloride to complete the quantitative transfer</li> <li>water bath temp 15-20 deg C above solvent boiling temp</li> <li>partially immerse concentrator tube in water and lower apparatus to complete concentration in 10-20 min</li> <li>sample removed from water bath when volume reaches 1 mL</li> <li>N bath temp 35 deg C</li> </ol>
QC - Spikes	Acid surrogate and spike components at 100 ug/mL; base/neutral surrogate and spike components at 50 ug/mL	Acid surrogate and spike components at 200 ug/mL; base/neutral surrogate and spike components at 100 ug/mL
QC - LCS	Acid surrogate and spike components at 100 ug/mL; base/neutral surrogate and spike components at 50 ug/mL	Acid surrogate and spike components at 200 ug/mL; base/neutral surrogate and spike components at 100 ug/mL

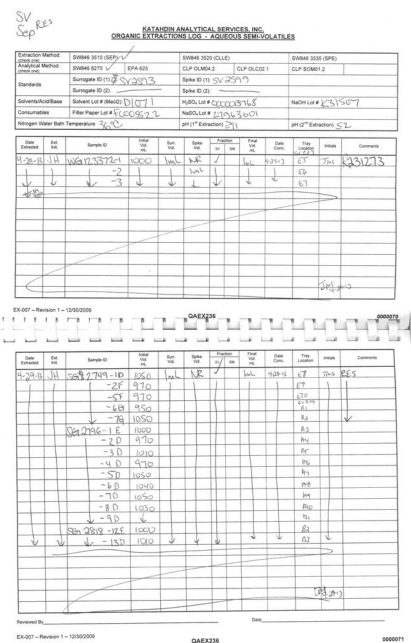
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# TITLE: PREPARATION OF AQUEOUS SAMPLES FOR EXTRACTABLE SEMIVOLATILE ANALYSIS

FIGURE 1

EXAMPLE OF SEMIVOLATILES LOGBOOK PAGE



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#### FIGURE 2

#### SOLVENT/REAGENT LOT CHECK LOGBOOK

GUMS

SOLVENT LOT CHECK

SOLVENT: A cotone

LOT#: D6818

DATE RECEIVED:

DATE CONCENTRATED: 4-8-13

CONCENTRATED BY: KF

PREP METHOD: JODM > /M

TRAY LOCATION: SLC2 (A9)

ANALYZED BY:

JUL

PASS/FAIL:

9

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# TITLE: PREPARATION OF AQUEOUS SAMPLES FOR EXTRACTABLE SEMIVOLATILE ANALYSIS

# FIGURE 3 LCS/MATRIX SPIKE COMPONENT LIST

BASE/NEUTRALS		
1-Methylnaphthalene	Bis (2-chloroethoxy) methane	
1,1-Biphenyl	Bis (2-chloroethyl) ether	
1,2,4-Trichlorobenzene	Bis (2-Chloroisopropyl) ether)	
1,2-Dichlorobenzene	Bis(2-Ethylhexyl)adipate	
1,3-Dichlorobenzene	Bis (2-ethylhexyl) phthalate	
1,4-Dichlorobenzene	Butylbenzyl phthalate	
1,4-Dioxane	Caprolactam	
2,4-Dinitrotoluene	Carbazole	
2,6-Dinitrotoluene	Chrysene	
2-Chloronaphthalene	Dibenz (a, h) anthracene	
2-Methylnaphthalene	Dibenzofuran	
2-Nitroaniline	Diethyl phthalate	
3,3'-Dichlorobenzidine	Diethyl adipate	
3-Nitroaniline	Dimethyl phthalate	
4-Bromophenylphenyl ether	Di-n-butylphthalate	
4-Chloroaniline	Di-n-octyl phthalate	
4-Chlorophenylphenyl ether	Fluoranthene	
4-Nitroaniline	Fluorene	
Acenaphthene	Hexachlorobenzene	
Acenaphthylene	Hexachlorobutadiene	
Acetophenone	Hexachlorocyclopentadiene	
Aniline	Hexachloroethane	
Anthracene	Indeno (1,2,3-cd) pyrene	
Atrazine	Isophorone	
Azobenzene	Naphthalene	
Benzaldehyde	Nitrobenzene	
Benzidine	N-Nitrosodimethylamine	
Benzo (a) Anthracene	N-Nitroso-di-n-propylamine	
Benzo (a) pyrene	N-Nitrosodiphenylamine	
Benzo (b) fluoranthene	Phenanthrene	
Benzo (ghi) perylene	p-toluidine	
Benzo (k) fluoranthene	Pyrene	
Benzyl alcohol	Pyridine	

ACIDS				
2, 3, 4, 6-Tetrachlorophenol	2-Chlorophenol	Benzoic acid		
2,4,5-Trichlorophenol	2-Methylphenol	Ethyl methanesulfonate		
2,4,6-Trichlorophenol	2-Nitrophenol	Methyl methanesulfonate		
2,4-Dichlorophenol	4,6-Dinitro-2-methylphenol	Pentachlorophenol		
2,4-Dimethylphenol	4-Chloro-3-methylphenol	Phenol		
2,4-Dinitrophenol	4-Methylphenol			
2,6-Dichlorophenol	4-Nitrophenol			

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# TITLE: PREPARATION OF AQUEOUS SAMPLES FOR EXTRACTABLE SEMIVOLATILE ANALYSIS

# FIGURE 4 APPENDIX IX LCS/MATRIX SPIKE COMPONENT LIST

	_
1,2,4,5-Tetrachlorobenzene	Hexachloropropene
1,3,5-Trinitrobenzene	Isodrin
1,4-Naphthoquinone	Isosafrole
1-Chloronaphthalene	Kepone
1-Naphthylamine	m-Dinitrobenzene
2,4-D	Methapyrilene
2-Acetyl aminofluorene	Methyl parathion
2-Naphthylamine	n-Nitrosodiethylamine
2-Picoline	n-Nitrosodi-n-butylamine
3,3-Dimethylbenzidine	n-Nitrosomethylethylamine
3-Methylcholanthrene	n-Nitrosomorpholine
4-Aminobiphenyl	n-Nitrosopyrrolidine
4-Nitroquinoline-1-oxide	n-Nitrotrosopiperidine
5-Nitro-o-toluidine	O,O,O-Triethyl phosphorothioate
7,12-Dimethylbenz(a)anthracene	o-Toluidine
a,a-Dimethylphenethylamine	Parathion
Acetophenone	p-Dimethylaminoazobenzene
Aramite	Pentachlorobenzene
Chlorobenzilate	Pentachloronitriobenzene
Diallate	Phenacetin
Dibenz(a,j)acridine	Phorate
Dimethoate	p-Phenylenediamine
Dinoseb	Pronamide
Diphenylamine	Safrole
Disulfoton	Silvex (2,4,5-TP)
Famphur	Sulfotep
Hexachlorophene	Thionazin

# KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-512 Revision History Cover Page Page 1

TITLE:		TION OF SEDIMENT/SOIL SAMPLES BY SESUBSEQUENT EXTRACTABLE SEMI-VOLA		
Prepared I	Ву:	Mike Thomas	Date:	09/96
Approved	Ву:			
Group Sup	pervisor:	Michael F. Thomas	Date:	11/15/00
Operations	s Manager:	CBento	Date:	10/25/00
QA Officer	<del>.</del> .	Deborah J. Nadeau	Date:	10.24.00
General M	anager:	Dermer P. Kufuss	Date:	11/16/00

Revision History:

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
01	Minor changes throughout. Clarifications to procedure Section.	Dn	10-24-00	10/24/00
02	Addition of Compounds to Figure 2.	Dr	3.28.02	3.28.02
03	Definitions added to section 1.1. Wording was added or changed to clarify sections 4,5,6,7,8+9. Hinor changes throughout. New figures.	HRC	11.08.04	11.08.04
04	Updated sect. 5.0 with current spike solutions prep. Removed section on medium level soil extraction, Replaced Figure 3 and 4 with current LCS/MS spike components, Minor corrections to sect. 1.3, 4.24,60 and 7.12. Updated logbook	LAD	04/06	04/06
<u>ک</u> و	Many Changes made throughout, including but not limited to, was te information, updated spikes and surrogates, added SIM LCS/D and MS/D information, updated Table 1. Please refer to the QAM SOP change form filed w/ SOP in QA for a detailed list of		09/07	69/07

SOP Number: CA-512 Revision History Cover Page (cont.) Page 2

TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
06	Updated logiocok example. Added addipute compounds to fig. J. Added necessity of recording lot numbers of consum ables in logiocok. Added to record the temperature of the nitrogen evaporation water booth.	CAN	80110	80/10
07	Added requirement to add spike before NaSOV. Changed N2 waterboth temperature from <39°C to 230°C femoved respirator reference. Added KAEHS manual. Added KASSOP CA-103, reference for a datitional Subsempling information.	LAN	09(09	oəloq
08	Removed targeting sample weights. Added KAT. SOP SD-902 reference. Updated logbook page example. Added GPC cleanup is required for all samples. Removed decenting samples	, LAD	08110	08/10
09	Minor modifications made to sections 5 & 7 to reflect current practices. Updated Sections, to include LOD/LOQ requirements, Changed 7.6°. T.7 to add surrogate and spikes after sodium so take is added. Updated references in Section 10.	LA D	04/12	04/12
	, 0			

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TITLE:	PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS
	acknowledge receipt of this standard operating procedure by signing and dating both of the provided. Return the bottom half of this sheet to the QA Department.
SEDIME	vledge receipt of copy of document SOP CA-512-09, titled PREPARATION OF INT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT CTABLE SEMI-VOLATILES ANALYSIS.
Recipier	nt:Date:
	DIN ANALYTICAL SERVICES, INC.
I acknow <b>SEDIME</b>	ARD OPERATING PROCEDURE  vledge receipt of copy of document SOP CA-512-09, titled PREPARATION OF  ENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT  CTABLE SEMI-VOLATILES ANALYSIS.
Recipier	nt: Date:

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

#### 1.0 SCOPE AND APPLICATION

The purpose of this SOP is to describe the procedures and requirements for the preparation of solid samples for analysis of extractable semivolatile organic compounds. This SOP is specifically applicable to EPA Method 3550B in accordance with SW-846 Method 8270, current revision.

#### 1.1 Definitions

METHOD BLANK (laboratory reagent blank): An artificial sample designed to determine if method analytes or other interferences are present in the laboratory environment, the reagents, or the apparatus. For aqueous samples, reagent water is used as a blank matrix; for soil samples, baked organic-free sand is used as a blank matrix. The blank is taken through the appropriate steps of the process.

LABORATORY CONTROL SAMPLE (LCS): A blank that has been spiked with the analyte(s) from an independent source, and is analyzed exactly like a sample. Its purpose is to determine whether the methodology is in control, and whether the laboratory is capable of making accurate and precise measurements. The matrix used should be phase matched with the samples and well characterized.

MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD): Predetermined quantities of stock solutions of certain analytes are added to a sample matrix prior to sample extraction and analysis. Samples are split into duplicates, spiked and analyzed. Percent recoveries are calculated for each of the analytes detected. The relative percent difference between the samples is calculated and used to assess analytical precision.

SURROGATES: Organic compounds which are similar to analytes of interest in chemical composition, extraction and chromatography, but which are not normally found in environmental samples. These compounds are spiked into all blanks, standards, samples and spiked samples prior to analysis. Percent recoveries are calculated for each surrogate.

#### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of analysts experienced in the extraction of samples for semivolatile analysis. Each analyst must demonstrate and document their ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, current revision, "Personnel Training".

It is the responsibility of all Katahdin technical personnel involved in the extraction of samples for semivolatile analysis to read and understand this SOP, adhere to the procedures outlined, and to properly document their data in the appropriate lab

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### TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

notebook. Any deviations from the test or irregularities with the samples should also be recorded in the lab notebook and reported to the Department Manager or designated qualified data reviewer responsible for this data.

It is the responsibility of the Department Manager to oversee that members of their group follow this SOP, to ensure that their work is properly documented and to indicate periodic review of the associated logbooks.

#### 1.3 Safety

Users of this procedure must be cognizant of inherent laboratory hazards, proper disposal procedures for contaminated materials and appropriate segregation of hazardous wastes. The toxicity or carcinogenicity of each reagent used in this method has not been precisely defined; however, each chemical should be treated as a potential health hazard. A reference file of material safety data sheets is available to all personnel involved in the chemical analysis. Everyone involved with the procedure must be familiar with the MSDSs for all the materials used in this procedure.

Each qualified analyst or technician must be familiar with Katahdin Analytical Environmental Health and Safety Manual including the Katahdin Hazardous Waste Plan and must follow appropriate procedures. These include the use of appropriate personal protective equipment (PPE) such as safety glasses, gloves and lab coats when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location of all safety equipment. Each analyst shall receive a safety orientation from their Department Manager, or designee, appropriate for the job functions they will perform.

#### 1.4 Waste Disposal

Wastes generated during the preparation of samples must be disposed of in accordance with the procedures described in the current revision of the Katahdin Analytical Environmental Health and Safety Manual and SOP SD-903, "Sample Disposal," current revision. Expired standards are lab packed, placed in the Katahdin hazardous waste storage area, and disposed of in accordance with this SOP.

Any methylene chloride solvent waste generated during the rinsing of glassware etc. should be disposed of in the "D" waste stream satellite accumulation area nearest the point of generation. Acetone and methanol are considered flammable waste, and should be disposed of in the "O" waste stream satellite accumulation area nearest the point of generation. Post-extraction soil samples and sodium sulfate waste should be disposed of in the soil with organics "I" waste stream satellite accumulation area nearest the point of generation. Please refer to the current revision of SOP CA-107 for the location of satellite waste accumulation areas.

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TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

#### 2.0 SUMMARY OF METHOD

An approximate, greater than 30 gram portion of sediment/soil is mixed with anhydrous powdered sodium sulfate and extracted with 1:1 methylene chloride/acetone (v/v) using an ultrasonic probe. The methylene chloride extract is dried and concentrated to a volume of 1.0 mL.

#### 3.0 INTERFERENCES

Contaminants in solvents, reagents, glassware, and other sample processing hardware may cause method interferences such as discrete artifacts and/or elevated baselines in the total ion current profiles (TICPs). All of these materials routinely must be demonstrated to be free from interferences under the conditions of the analysis by running laboratory method blanks. Interferences caused by phthalate esters can pose a major problem in semivolatile organics analysis because many phthalates are also target analytes. Common flexible plastics contain varying amounts of phthalates that are easily extracted during laboratory operations, so cross-contamination of glassware frequently occurs when plastics are handled. Interferences from phthalates can best be minimized by avoiding the use of such plastics in the laboratory.

At no time may gloves that have not been tested for phthalates or gloves known to contain phthalates be used or stored in the organic extraction lab. Additionally, whenever possible plastic items in this lab must be replaced with metal or Teflon or other non-phthalate plastic substitute.

Special care should be taken to ensure clean glassware and apparatus are used, prerinsed with the appropriate solvent prior to use. Solvents should be analyzed prior to use to demonstrate that each lot is free of contaminants that may interfere with the analysis. Interferences coextracted from the samples will vary considerably from source to source. If analysis of an extracted sample is prevented due to inteferences, further cleanup of the sample extract may be needed to minimize interferences.

#### 4.0 APPARATUS AND MATERIALS

Prior to use, all glassware must be rinsed three times with methylene chloride. Brand names and catalog numbers are included below for illustration purposes only.

- 4.1 Syringe gas tight, 1.0 mL, solvent rinsed between each use.
- 4.2 Sonicator ultrasonic processor XL Misonix (or equivalent) equipped with dual titanium 3/4" horn extenders for extracting two samples at a time.
- 4.3 Powder funnels, 100 mm diameter, 35 mm stem

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# TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

- 4.4 Kuderna-Danish (KD) apparatus Concentrator tube 10 mL Evaporative flask - 500 mL Snyder column - 3-ball macro
- 4.5 Filter paper, 7.0 cm, Whatman #4
- 4.6 Vacuum filtration flask 500 mL Erlenmeyer
- 4.7 Buchner funnel, porcelain, Coors® with 85 mm plate diameter (or equivalent)
- 4.8 Beakers 400 mL
- 4.9 Boiling chips approximately 12 mesh, silicon carbide (carborundum or equivalent). Soxhlet extract overnight in methylene chloride.
- 4.10 Water bath eight position concentric ring bath, or equivalent, equipped with a calibrated thermometer. The bath should be used in a hood.
- 4.11 Balance capable of accurately weighing ± 0.01 g.
- 4.12 Vials and caps 1.8 mL with PTFE/silicone septa and 12 mL with Teflon-lined caps for extracts designated for GPC cleanup.
- 4.13 Filter paper, 18.5 cm, Fisherbrand or Whatman 114V (or equivalent)
- 4.14 Pasteur pipets disposable, 5 3/4 ".
- 4.15 Nitrogen evaporation apparatus.
- 4.16 Muffle oven capable of maintaining 400 °C for baking glass wool and organic-free sand.
- 4.17 Gel Permeation Chromatograph (GPC) J2 Scientific AccuPrep MPS<sup>™</sup> with internal UV detection

#### 5.0 REAGENTS

5.1 Sodium Sulfate - anhydrous powdered and granular crystals, reagent grade, certified by the manufacturer/vendor as purified heating to 400°C prior to receipt by the laboratory. (Jost Chemical anhydrous powder, catalog #2797 or equivalent, and Jost Chemical granular crystals, catalog #2796 or equivalent).

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# TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

- 5.2 Methylene chloride, methanol, and acetone pesticide residue analysis grade or equivalent. Methylene chloride and acetone are evaluated by lot prior to use by concentration of approximately 200 mL to 1.0 mL followed by GC/MS analysis. The lot numbers of all solvents used during an extraction must be recorded in the extraction logbook.
- 5.3 Organic-free sand, purified by baking at 400 °C. Method blanks serve as checks on the baked sand.
- 5.4 Base/Neutral and Acid (SVOA) Surrogate Spiking Solution Surrogate standards are added to all samples and calibration solutions. Prepare a surrogate standard spiking solution that contains the following compounds at the indicated concentrations in acetone.

Compound	Conc.
phenol- <sub>d6</sub>	100 ug/mL
2,4,6-tribromophenol	100 ug/mL
2-fluorophenol	100 ug/mL
nitrobenzene- <sub>d5</sub>	50 ug/mL
p-terphenyl- <sub>d14</sub>	50 ug/mL
2-fluorobiphenyl	50 ug/mL

Store the spiking solution at -10°C to -20°C in Teflon-sealed containers. These solutions must be replaced after six months or sooner if comparison with quality control check samples indicates a problem. If reanalysis of a method blank still indicates surrogates out of criteria, a new surrogate solution must be used.

5.5 SIM Surrogate Spiking Solution- Surrogate Standards are added to all samples and calibration solutions. Prepare a surrogate solution that contains the following compounds at a concentration of 2 ug/mL in acetone.

Compound	Conc. ug/mL
Fluorene-d10	2.0 ug/mL
2-Methylnaphthalene-d10	2.0 ug/mL
Pyrene-d10.	2.0 ug/mL
2,4-Dibromophenol	4.0 ug/mL
1,4-Dioxane-d8	2.0 ug/mL

Store the spiking solution at -10°C to -20°C in Teflon-sealed containers. These solutions must be replaced after six months, or sooner if comparison with quality control check samples indicates a problem. If reanalysis of a method blank still indicates surrogates out of criteria, a new surrogate solution must be used.

5.6 Base/Neutral and Acid (SVOA) Matrix Spike/Lab Control Sample Spiking Solution - Prepare a spiking solution in methanol that contains the compounds listed in Figure 2

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### TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

at a concentration of 50 ug/mL for base/neutrals and 100 ug/mL for acids. Store the spiking solution at -10°C to -20°C in Teflon-sealed containers. These solutions must be replaced after six months, or sooner if comparison with quality control check samples indicates a problem.

- 5.7 Base/Neutral and Acid (SVOA APPENDIX IX) Matrix Spike/Lab Control Sample Spiking Solution. Prepare a spiking solution in methanol that contains the compounds listed in Figure 3 at a concentration of 100 μg/mL for each compound. Store the spiking solution at -10°C to -20°C in Teflon-sealed containers. These solutions must be replaced after six months, or sooner if comparison with quality control check samples indicates a problem
- 5.8 Base/Neutral and Acid (SIM) Matrix Spike/ Lab Control Sample Spike Solution for SIM-SVOA. Prepare a spiking solution in methanol that contains the compounds listed in Figure 2 at a concentration of 2.0 ug/mL for base/neutral and 4.0 ug/mL for acid. Take out 1.0 mL of Base/Neutral and Acid Matrix Spike/Lab Control Spiking Solution for SVOA and dilute it to 25.0 mL in methanol. Store the solution Spiking at 10°C to -20°C in Teflon-sealed containers. These solutions must be replaced after six months, or sooner if comparison with quality control check samples indicates a problem.

### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

Sediment/soil samples must be collected in a soil jar and must be maintained at 4°C (±2°C).

Holding time for extraction of sediment/soil samples for Method 3550 is 14 days from date of sample collection, although the analyst should be aware that actual holding times employed may be project/program specific.

Store all extracts at 4°C (±2°C) in the dark in labeled Teflon-sealed containers. See SOP SD-902, "Sample Receipt and Internal Control," current revision, for storage areas and temperature maintenance procedures.

#### 7.0 PROCEDURES

The following information must be recorded in the extraction logbook.

- Extraction method
- Surrogate and spike IDs
- Lot numbers of all solvents, acids and bases, sodium sulfate, filter paper
- Nitrogen evaporation water bath temperature
- pH if applicable

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### TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

- Sonicator horns tuned
- Extraction and Concentration dates
- Extraction and Concentration analyst
- Sample ID or QC sample ID
- Initial and final volumes or weight
- Surrogate and spike amounts
- Final extract tray location
- Any comments regarding the sample extraction (ie. Emulsion)

All solid samples should be cleaned using gel permeation chromatography (GPC) to reduce matrix interferences.

The organic department manager should be consulted to determine if a particular sample should be subjected to further cleanup procedures; the decision should consider sample history, sample appearance, and project/client needs.

Samples need to be "swiped" out when removing and "swiped" in when replacing samples in storage locations to maintain the internal chain of custody. Refer to Katahdin SOP SD-902, Sample Receipt and Internal Control, current revision, for the proper procedure for removal and return samples. Fill out the sample preparation/extraction log with the necessary information before starting the extraction. Prerinse all glassware three times with methylene chloride.

- 7.1 Do not decant any water layer on a sediment sample. Mix with a stainless steel spatula to ensure homogeneity of the sample. If the sample container is full to the extent that stirring the sample is impractical, try to remove the "best representative" aliquot from the jar based on color, particle size, moisture, etc. Remove any foreign objects such as sticks, leaves, and rocks, and note actions taken in the appropriate extraction logbook. Please refer to the current revision of Katahdin Analytical Services SOP CA-108, "Basic Laboratory Technique", for more detailed guidance on subsampling to ensure reproducibility.
- 7.2 The following steps should be performed <u>rapidly to avoid loss of the more volatile extractable</u>. Weigh out an approximate, greater than 30g portion of sample into a labeled 400-mL beaker. Record sample weight to the nearest 0.01 g in appropriate extraction logbook. Add between 30 g and 60 g of anhydrous powdered sodium sulfate as required for producing a "free-flowing" mixture. The amount of sodium sulfate added will depend upon the moisture content of the sample (e.g., low moisture content will require less sodium sulfate). Mix well with a spatula. Keep the spatula in the sample beaker and <u>cover</u> the beaker with aluminum foil.
- 7.3 A method blank must be prepared with each extraction batch, not to exceed 20 client samples. To prepare a method blank, weigh out one, greater than 30 g portion of purified sand in a labeled 400 mL beaker. Refer to sections 7.6 and 7.7

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### TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

for spike and surrogate addition instructions. Add 60 g sodium sulfate and mix well. Although a "free-flowing" mixture can be achieved with less than 60 g sodium sulfate, the method blank must contain 60 g in order to evaluate the sodium sulfate as a potential source of contamination.

- 7.4 A laboratory control sample (LCS) must be prepared with each extraction batch, not to exceed 20 client samples. To prepare LCS, weigh out one 30 g portion of purified sand in a labeled 400 mL beaker. Refer to sections 7.6 and 7.7 for spike and surrogate addition instructions. Add 30 g sodium sulfate and mix well. If combined SIM-SVOA analysis is requested, a separate LCS must be prepared for each analysis. If an MS/MSD pair is not extracted on a particular day, an LCS/LCSD pair may be required in order to meet client-specific or program-specific requirements. This information will be disseminated from the project manager or Department Manager.
- 7.5 A matrix spike/matrix spike duplicate (MS/MSD) set must be prepared for every 20 samples. To prepare MS/MSD, weigh out two approximate, greater than 30 g portions of the sample designated for MS/MSD into each of two labeled 400 mL beakers. Record sample weights to nearest 0.01 g in appropriate extraction logbook. Refer to sections 7.6 and 7.7 for spike and surrogate addition instructions. Add 30 60 g sodium sulfate to each to produce a free-flowing mixture, and mix well. If combined SIM-SVOA analysis is requested, a separate MS/MSD must be prepared for each analysis.
- 7.6 To all samples, method blank, LCS/LCSD, and MS/MSD add 1.0 mL of the appropriate base/neutral and acid surrogate spiking solution listed below using the pre-rinsed 1.0 mL gas tight syringe. The surrogate spike should be added **after** the addition of the sodium sulfate. Record surrogate spike volume and identification code in extraction logbook. Thoroughly rinse syringe with solvent prior to using it for another spiking solution.
  - 7.6.1 If the request is for SVOA or SVOA Appendix IX, use the SVOA surrogate solution (sect. 5.4).
  - 7.6.2 If the request is for SIM, use the SIM surrogate solution (sect. 5.5).
  - 7.6.3 If the request is for SIM-SVOA, use both the SIM and SVOA surrogate solutions. NOTE: Separate LCS/LCSD and/or MS/MSD are needed for each analysis and should be spiked with the appropriate surrogate.
- 7.7 To the LCS/LCSD and the MS/MSD add 1.0 mL of the appropriate base/neutral and acid (SVOA) matrix spike/LCS spiking solution listed below using a 1.0 mL gas tight syringe. The LCS/MS spike should be added **after** the addition of the sodium sulfate. Record the matrix spike/LCS spiking solution volume and identification

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code in extraction logbook. Thoroughly rinse the syringe with solvent when spiking is completed.

- 7.7.1 If the request is for SVOA, add 1 mL of SVOA Spiking Solution (sect 5.6).
- 7.7.2 If the request is for SIM, add 1 mL SIM Spiking solution (sect 5.8).
- 7.7.3 If the request is for SVOA and SIM, add 1mL of SVOA Spiking Solution and 1 mL SIM Spiking solution (sect 5.6 and 5.8).
- 7.7.4 If the request is for SVOA Appendix IX, add 1mL of SVOA Spiking Solution and 1 mL of SVOA Appendix IX Spiking solution (sect 5.6 and 5.7).
- 7.8 To assure optimum operation and maximum energy output, the sonicators <u>must</u> be tuned daily prior to extracting samples. The following tuning procedure must be performed with the sonicator probes vibrating in air.
  - 7.8.1 Turn OUTPUT CONTROL knob counter-clockwise to zero. This automatically switches the duty cycle to continuous mode.
  - 7.8.2 Press and hold down the power switch to on.
  - 7.8.3 Press and hold down the TUNE switch. Check if the counter is less or equal to 20%; otherwise, rotate the Tuning Knob (tuning button) clockwise until a reading of 20% ress is obtained.
  - 7.8.4 Release the TUNE switch.
  - 7.8.5 Turn OUTPUT CONTROL KNOB counter-clockwise to 50 and the power switch off.
  - 7.8.6 Confirm that the sonicators were tuned by recording the date and/or percent in the extractions logbook.
- 7.9 Prior to extracting any samples, ensure that the sonicator probes are decontaminated by rinsing three times with a methylene chloride wash bottle. Collect the waste in a waste beaker. It may sometimes be necessary to wipe the upper part of each probe with a methylene chloride dampened KimWipe. Repeat this decontamination step between each sample on each probe.
- 7.10 To the mixed and spiked blank and LCS, add approximately 100 mL of the 1:1 methylene chloride/acetone (V/V) solution and proceed with steps 7.11 through 7.14. Record the lot numbers of the solvents in the extraction logbook.

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- 7.11 It may be necessary at this time to stir the sample/sodium sulfate mixture with the spatula to loosen up the mixture prior to extracting. Rinse the spatula with methylene chloride and collect the rinsing into a correspondent beaker. Position the beaker in the ultrasonic cell disruptor so that the bottom surface of the tip of the 3/4 inch disruptor horn is about halfway below the surface of the solvent and above the sediment layer.
- 7.12 Sonicate for 3 minutes with the output control knob set at 10, and mode switch on "pulsed" and % duty cycle knob set at 50%. While the mixture is sonicating, one should be able to see all, or most of the material, moving in the beaker under the influence of the energized probes. If not, stir the mixture again.
- 7.13 Prepare a filter flask fitted with a Buchner funnel. The Buchner funnel should contain a 7.0 cm Whatman #4 filter. Prerinse the flask, funnel and filter with methylene chloride and discard rinsings into solvent waste container. Decant extract into the filter flask and Buchner funnel. A vacuum pump may be used to facilitate filtration or the extract may be gravity filtered. The lot number of the filter paper must be written ti the extraction logbook.
- 7.14 Repeat the extraction two more times (sec 7.11 7.14) using approximately 100 mL portions of 1:1 methylene chloride: acetone. Before each extraction, make certain that the sodium sulfate is still free-flowing and not a consolidated mass. As required, break up large lumps with the spatula. Decant the extraction solvent into the Buchner funnel after each sonication. On the final sonication, pour the entire sample contents into the Buchner funnel and rinse thoroughly with methylene chloride to complete the quantitative transfer of the extract. Use the vacuum pump to pull all the extract into the flask

#### CONCENTRATION OF LOW LEVEL EXTRACTS

- 7.15 Rinse the K-D glassware (flask, concentration tube, and Snyder column, including the ground glass joints on the flask and columns) three times with methylene chloride before assembling. Add two boiling chips to the K-D. Insert 18.5 cm filter papers into short stem powder funnels and add ~ 2 inches of sodium sulfate crystals. Place the assembled K-D's under the funnels. The lot number of the filter paper must be written in the extraction logbook.
- 7.16 Transfer the extracts to the K-D concentrator setups through the sodium sulfate in the funnels. This is the drying step, which is required to remove residual water from the extracts. Any large water layers must be removed by other means, prior to pouring through the sodium sulfate. After pouring all of the extract volume through the sodium sulfate, rinse the extract flask three times with ~ 2 3 mls of methylene chloride. Add the rinsings through the sodium sulfate to complete a quantitative

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### TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

transfer. Rinse the sodium sulfate with ~ 15 mls of methylene chloride and allow to drain.

- 7.17 If samples are to be GPC'd, refer to the current revision of Katahdin SOP CA-513, Extract Cleanup Using Gel Permeation Chromatography, for appropriate concentrating procedures.
- 7.18 If samples are not to be GPC'd, follow Steps 7.19 through 7.24 to concentrate extracts to final volume of 1 mL. Otherwise proceed to GPC cleanup procedure as described in the current revision of Katahdin SOP CA-513.
- 7.19 Transfer the labels from the extract flasks to the K-Ds. Remove the funnel and attach a 3-ball macro Snyder column. Pre-wet the Snyder column with 1 mL of methylene chloride.
- 7.20 Place the K-D in a hot water bath (75-85°C). Gently swirl K-D in the water until boiling begins. At the proper distillation rate, the Snyder balls should chatter but the chambers should not flood with condensed solvent. The K-D should be kept in a vertical orientation while on the bath. When the apparent volume in the concentrator tube reaches ≈ 6 mL, remove the K-D from the water bath. Do not allow the evaporator to go dry. If the sample extract does go dry, re-extraction must occur immediately. Allow the K-D to cool for 10 minutes. Rinse the Snyder column lower joint with ≈ 1 mL of methylene chloride. Remove the Snyder column. Wipe off any water from the neck above the lower joint of the flask. Separate the K-D flask from the concentrator tube, rinsing the ground glass joint with ≈ 1 mL methylene chloride.
- 7.21 Reduce the extract in the concentrator tube to approximately 1 mL using the nitrogen blow-down apparatus. The bath temperature must be < 39°C. Turn the gas to 3 psi. Be careful not to splash the extract out of the tube. <u>During concentration on the N-evap</u>, the internal wall of the concentrator tube and the N-evap sparging pipet must be rinsed down at least once or twice with ≈1 mL of methylene chloride. The solvent level in the concentrator tube must be positioned below the level of the water bath as much as possible to prevent water from condensing into the sample extract. As the extract volume is reduced, lower the N₂ sparging pipet closer to the surface of the extract to expedite the concentration. Record the temperature of the water in the nitrogen evaporation water bath in the logbook also note any problems or extract losses, if they occur, in the extractions logbook.
- 7.22 When the apparent volume reaches slightly less than 1 mL, remove the concentrator tube and allow it to cool.

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- 7.23 Complete the quantitative transfer of the extract to a 1.8 mL vial by using methylene chloride. Adjust the volume of the methylene chloride extract to 1.0 mL using the 1.8 mL reference vial for volume comparison.
- 7.24 Label the vial with lab sample number, extraction date, matrix and analysis. Store extract vials at a temperature of 4 ± 2 °C until ready for analysis. Indicate in the extraction logbook the box number and "tray location" of the individual extract vials.

#### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

A method blank must be extracted for each and every item listed below:

- Each sample matrix
- Each extraction method or level
- Every extraction batch of twenty or fewer samples

A laboratory control sample (LCS) is required for <u>each</u> and <u>every</u> item listed below:

- Each sample matrix
- Each extraction method or level
- Every extraction batch of twenty or fewer samples

Refer to the current revision of the applicable Katahdin SOP for analysis of Semivolatile Organics for quality control acceptance criteria.

Each extractions analyst must demonstrate proficiency in performing the extractions that prepare samples for analysis. Demonstration consists of preparation (extraction), by the analyst, of at least four aliquots which are then analyzed according to the analytical method in question. These QC samples must meet all quality control acceptance limits. Demonstration must be documented by use of a form which summarizes the results of the analysis of these aliquots, calculated percent recoveries, and standard deviation. Demonstration of proficiency must be done one time per analyst initially and then annually thereafter. Refer to SOPs QA-805 and QA-807, current revision.

If, upon analysis of the extracted samples, it is discovered that quality control acceptance criteria have not been met, all associated samples must be evaluated against all the QC. In some cases data may be reported, perhaps with narration, while in other cases, other corrective action may be taken. The corrective actions may include re-extraction of the samples associated with the quality control sample that did not meet acceptance criteria, or may include making new reagents and standards if the standardization is suspect. These decisions are based on holding time considerations, client and project specific Data Quality Objectives and on review of chromatograms. The supervisor, Operations Manager, and/or Quality Assurance Officer may be consulted to evaluate data. Some samples may not be able

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to be reanalyzed within hold time. In these cases "qualified" data with narration may be advisable after consultation with the client.

Much of the work performed at the lab is analyzed in accordance with specific QC requirements spelled out in a project specific Quality Assurance Project Plan (QAPP) or in a program specific Quality Systems Manual (QSM). The reporting limits, acceptance criteria and/or corrective actions may be different than those specified in this SOP. In these cases the appropriate information will be communicated to the Department Manager and/or senior chemists before initiation of the analyses so that specific product codes can be produced for the project. In addition, the work order notes for each project will describe the specific QAPP or QSM to be followed.

#### 9.0 METHOD PERFORMANCE

The method detection limit (MDL) is defined as the minimum concentration of a substance that can be measured and reported with 99% confidence that the value is above zero. The MDLs shall be determined and verified one time per type of instrument unless otherwise required by the method.

A Limit of Detection (LOD) is an estimate of the minimum amount of a substance that an analytical process can reliably detect. An LOD is analyte and matrix specific and may be laboratory-dependent. LODs must be determined for all parameters for which the laboratory is accredited under the DoD Environmental Laboratory Accreditation Program. LOD's must be verified for every preparation and analytical method combination and on every applicable instrument on a quarterly basis.

The Limit of Detection (LOQ) is the minimum levels, concentrations, or quantities of a target variable (e.g., target analyte) that can be reported with a specified degree of confidence. The LOQ shall be set at the lowest point in the calibration curve for all analyses utilizing an initial calibration. LOQ's must be verified quarterly for every preparation and analytical method combination and on every applicable instrument on a quarterly basis for all parameters included in the DoD Scope of Accreditation. The LOQ must be verified at least once annually if the analysis is not included in the DoD Scope of Accreditation.

MDLs are filed with the Organic Department Manager and then with the QAO. LOD and LOQ verifications are filed with the QAO

Refer to the current revision of Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, for procedures on determining the MDL.

Refer to the applicable analytical SOP for other method performance parameters and requirements.

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TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

### 10.0 APPLICABLE DOCUMENTS/REFERENCES

Test Methods for Evaluating Solid Waste-Physical/Chemical Methods, Method 3550C, USEPA SW-846, Third Edition, Update IV, February 2007.

The NELAC Institute, Laboratory Accreditation Standards, Volume 1, Management and Technical Requirements for Laboratories Performing Environmental Analysis, 10/06/2010.

Department of Defense Quality Systems Manual for Environmental Laboratories (DOD QSM), Current Version.

Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, current revision.

#### LIST OF TABLES AND FIGURES

Table 1 Summary of Method Modifications

Figure 1 Example of Logbook Page

Figure 2 LCS/Matrix Spike Component List

Figure 3 Appendix IX LCS/Matrix Spike Component List

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### TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

# TABLE 1 SUMMARY OF METHOD MODIFICATIONS

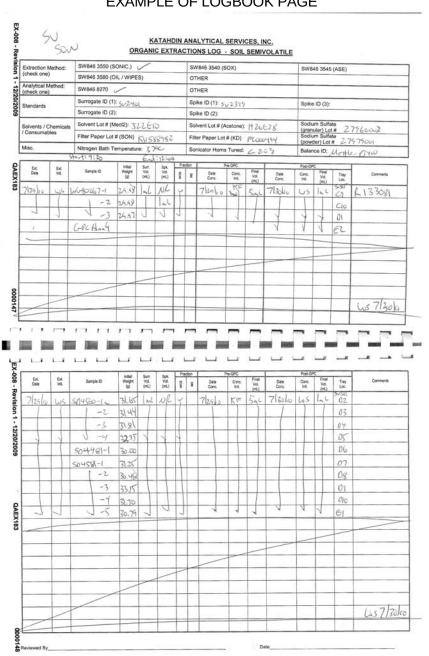
TOPIC	KATAHDIN SOP CA-512-09	METHOD 3550, current revision
Apparatus/Materials	1) short stem funnels	1) drying columns
Reagents		
Sample preservation/ handling		
Procedures	<ol> <li>extract dried using Na<sub>2</sub>SO<sub>4</sub> in short stem funnels</li> <li>place sonicator horns ½ way between the surface of the solvent and the sediment layer</li> <li>no apparatus height specification for concentration on water bath</li> <li>water bath at 75-85 deg C</li> <li>sample removed from water bath when volume reaches ~6 mL</li> </ol>	<ol> <li>extract dried using Na<sub>2</sub>SO<sub>4</sub> in drying columns</li> <li>place sonicator horns ½ inch below the solvent surface but above sediment layer</li> <li>partially immerse concentrator tube in water and lower apparatus to complete concentration in 10-15min</li> <li>water bath at 80-90 deg C</li> <li>sample removed from water bath when volume reaches 1-2 mL</li> </ol>
QC - Spikes	Acid surrogate and spike components at 100 ug/mL; base/neutral surrogate and spike components at 50 ug/mL	Acid surrogate and spike components at 200 ug/mL; base/neutral surrogate and spike components at 100 ug/mL
QC - LCS	Acid surrogate and spike components at 100 ug/mL; base/neutral surrogate and spike components at 50 ug/mL	Acid surrogate and spike components at 200 ug/mL; base/neutral surrogate and spike components at 100 ug/mL
QC - Accuracy/Precision		
QC - MDL		

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### TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

FIGURE 1

EXAMPLE OF LOGBOOK PAGE



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### TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

# FIGURE 2 LCS/MATRIX SPIKE COMPONENT LIST

BASE/NEUTRALS		
1-Methylnaphthalene	Bis (2-chloroethoxy) methane	
1,1-Biphenyl	Bis (2-chloroethyl) ether	
1,2,4-Trichlorobenzene	Bis (2-Chloroisopropyl) ether)	
1,2-Dichlorobenzene	Bis (2-ethylhexyl) adipate	
1,3-Dichlorobenzene	Bis (2-ethylhexyl) phthalate	
1,4-Dichlorobenzene	Butylbenzyl phthalate	
1,4-Dioxane	Caprolactam	
2,4-Dinitrotoluene	Carbazole	
2,6-Dinitrotoluene	Chrysene	
2-Chloronaphthalene	Dibenz (a, h) anthracene	
2-Methylnaphthalene	Dibenzofuran	
2-Nitroaniline	Diethyl adipate	
3,3'-Dichlorobenzidine	Diethyl phthalate	
3-Nitroaniline	Dimethyl phthalate	
4-Bromophenylphenyl ether	Di-n-butylphthalate	
4-Chloroaniline	Di-n-octyl phthalate	
4-Chlorophenylphenyl ether	Fluoranthene	
4-Nitroaniline	Fluorene	
Acenaphthene	Hexachlorobenzene	
Acenaphthylene	Hexachlorobutadiene	
Acetophenone	Hexachlorocyclopentadiene	
Aniline	Hexachloroethane	
Anthracene	Indeno (1,2,3-cd) pyrene	
Atrazine	Isophorone	
Azobenzene	Naphthalene	
Benzaldehyde	Nitrobenzene	
Benzidine	N-Nitrosodimethylamine	
Benzo (a) Anthracene	N-Nitroso-di-n-propylamine	
Benzo (a) pyrene	N-Nitrosodiphenylamine	
Benzo (b) fluoranthene	Phenanthrene	
Benzo (ghi) perylene	p-toluidine	
Benzo (k) fluoranthene	Pyrene	
Benzyl alcohol	Pyridine	

ACIDS				
2, 3, 4, 6-Tetrachlorophenol	2-Chlorophenol	Benzoic acid		
2,4,5-Trichlorophenol	2-Methylphenol	Ethyl methanesulfonate		
2,4,6-Trichlorophenol	2-Nitrophenol	Methyl methanesulfonate		
2,4-Dichlorophenol	4,6-Dinitro-2-methylphenol	Pentachlorophenol		
2,4-Dimethylphenol	4-Chloro-3-methylphenol	Phenol		
2,4-Dinitrophenol	4-Methylphenol			
2,6-Dichlorophenol	4-Nitrophenol			

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### TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SONICATION USING METHOD 3550 FOR SUBSEQUENT EXTRACTABLE SEMI-VOLATILES ANALYSIS

# FIGURE 3 APPENDIX IX LCS/MATRIX SPIKE COMPONENT LIST

1,2,4,5-Tetrachlorobenzene	Hexachloropropene
1,3,5-Trinitrobenzene	Isodrin
1,4-Naphthoquinone	Isosafrole
1-Chloronaphthalene	Kepone
1-Naphthylamine	m-Dinitrobenzene
2,4-D	Methapyrilene
2-Acetyl aminofluorene	Methyl parathion
2-Naphthylamine	n-Nitrosodiethylamine
2-Picoline	n-Nitrosodi-n-butylamine
3,3-Dimethylbenzidine	n-Nitrosomethylethylamine
3-Methylcholanthrene	n-Nitrosomorpholine
4-Aminobiphenyl	n-Nitrosopyrrolidine
4-Nitroquinoline-1-oxide	n-Nitrotrosopiperidine
5-Nitro-o-toluidine	O,O,O-Triethyl phosphorothioate
7,12-Dimethylbenz(a)anthracene	o-Toluidine
a,a-Dimethylphenethylamine	Parathion
Acetophenone	p-Dimethylaminoazobenzene
Aramite	Pentachlorobenzene
Chlorobenzilate	Pentachloronitriobenzene
Diallate	Phenacetin
Dibenz(a,j)acridine	Phorate
Dimethoate	p-Phenylenediamine
Dinoseb	Pronamide
Diphenylamine	Safrole
Disulfoton	Silvex (2,4,5-TP)
Famphur	Sulfotep
Hexachlorophene	Thionazin

# ADDENDUM SOP NO CHANGE FORM

### KATAHDIN ANALYTICAL SERVICES, INC. SOP "REVIEW WITH NO CHANGES" FORM

Name of Person Reviewing SOP: Tessica Spear	h-Wildes
Review Date: 3-1513	
SOP Number: OA-5/2	
SOP Title: Preparation of Seliment	( Soil Samples by Sonication
sop Title: Preparation of Sediment Using method 3550 For Semi	i-volatives
THE ABOVE REFERENCED SOP HAS BEEN REVIEWED ANALYST OR SUPERVISOR. NO CHANGES ARE REQU	BY A QUALIFIED AND TRAINED
Department Supervisor Signature:	Date:
Hitim J	3-19-13
QAO Signature:	Date:
Liseio Dimno	031913

SOP Number: CA-213 Revision History Cover Page Page 1

TITLE:	ANALYSIS OF SEMIVOLATILE ORGANIC COMPOUNDS BY: SW 846 METHOD 8270
	- Modified for Selected Ion Monitoring (SIM)

Prepared By:	GC/MS Department	Date:	6/98
Approved By:	1		
Group Supervisor:	A Halog	Date:	020101
Operations Manager:	Joh C. Benton	Date:	1/31/01
QA Officer:	Quetorah J. Nadeau	Date:	1.31.01
General Manager:	Dunger J. Lukan	Date:	1/01/01
Revision History:			

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
01 8270C Mod	Format changes added pollution prevention added instrument and other calibration options. Other minor changes to sections 7,8 , OATable.	Dh	1:31:01	131.01
02 8270C	Many changes in formatting. Some additions to section & + Table 1 to comply with Navy.	Ðn	09:3004	09:30:04
03 8270C	Sect. 7.2: Removed "K" Instrument : added "R" instrument. Added Pentafluorophenol sur. to Tables 3, 5 and Sect. 8.2. Removed all references to TIC's.	LAO	04/06	04/06
	Sect. 8.2 - changed 5 to 4 and removed pentachlorophenol. Table 3 and 5 - removed pentachlorophenol. Changed linear regression correlation coefficient criteria. Added MISOP reference. Added LCS exceedance criteria. Added ICV requirementand criteria. Added RT vindow procedure.	LAV	06/07	06/07
05 8270C	Added "G" instrument, Removed "X" instrument Edited Section 7.5.1-initial cal table	UAN	02/08	02/08

### KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-213 Revision History Cover Page – Cont.

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SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
06	Section S.3.2.3- Added cerlibration Mix B. Section 7.5.1- Edited to address differt SIM compounds may need to be calibrated at different levels depending on the compound and project requirements.	LAD	04/09	04/09
07	Changes made for compliance with DoDasm version 4.1	LAD	08109	08/09
08	Updated Standard Prep. Added Compounds to Table 3 and 5. Updated references. Added DoDQSMQC requirements Table.	UAO	04/10	04/10
09	Sect. 7.4- Added additional tune information. Sect. 7.6- Added 100 w minimum extract vol. & I we IS is added for each 100 ve cliquest. Sect. 7.5.4- Added RRT Information. Sect. 9.0- Added MDL, LOD and LOO information. Table 4- Added 1,4-Dioxane-de Survo	LAD	oeln	05 (n
10	Sect. 7-Changed sample volume from Ind to 2nd. Sect. 8- Added 10% or vie for non-DoD clients. Sect. 9-Added MOL LOD and LOQ information. Sect. 10-Added and updated references. Updated Figure 1. Added Addedendum 1- LOW level 1. 4-Dioxane analysis	LAO	05/12	os liz
Production Production	Sect. I and 7. Removed Quickform reporting and added KIMS. Sect. 8 and Table 1. Added the surregate i.4-Dioxand 18. Throughout - Fixed typos and made minor changes.	LAD	03/13	03/13

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	acknowledge receipt of this standard operating procedure by signing and dating both aces provided. Return the bottom half of this sheet to the QA Department.	ı of
<b>SEMIVO</b>	owledge receipt of copy of document SOP CA-213-11, titled "ANALYSIS OF OLATILE ORGANIC COMPOUNDS BY METHOD 8270 – Modified for Selected Ion oring (SIM)".	
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#### 1.0 SCOPE AND APPLICATION

The purpose of this SOP is to describe the procedures utilized by Katahdin Analytical Services, Inc. laboratory personnel to prepare and analyze water and soil sample extracts for semivolatile organics by EPA SW-846 Method 8270, current revision, modified for selected ion monitoring.

In order to maintain consistency in data quality, this SOP consolidates all aspects of the analyses in one working document, to be revised as necessary.

#### 1.1 Definitions

ANALYTICAL BATCH: 20 or fewer samples which are analyzed together with the same method sequence and the same lots of reagents and with the manipulations common to each sample within the same time period or in continuous sequential time periods.

METHOD BLANK (laboratory reagent blank): An artificial sample designed to determine if method analytes or other interferences are present in the laboratory environment, the reagents, or the apparatus. For aqueous samples, reagent water is used as a blank matrix; for soil samples, baked organic-free sand is used as a blank matrix. The blank is taken through the appropriate steps of the process.

CALIBRATION CHECK: Verification of the ratio of instrument response to analyte amount, a calibration check is done by analyzing for analyte standards in an appropriate solvent. Calibration check solutions are made from a stock solution that is different from the stock used to prepare standards.

CALIBRATION STANDARD (WORKING STANDARD): A solution prepared from the stock standard solution that is used to calibrate the instrument response with respect to analyte concentration.

LABORATORY CONTROL SAMPLE (LCS): A blank that has been spiked with the analyte(s) from an independent source and is analyzed exactly like a sample. Its purpose is to determine whether the methodology is in control, and whether the laboratory is capable of making accurate and precise measurements. The matrix used should be phase matched with the samples and well characterized.

MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD): Predetermined quantities of stock solutions of certain analytes are added to a sample matrix prior to sample extraction and analysis. Samples are split into duplicates, spiked and analyzed. Percent recoveries are calculated for each of the analytes detected. The relative percent difference between the samples is calculated and used to assess analytical precision.

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STANDARD CURVE (CALIBRATION CURVE): A curve that plots concentration of known analyte standard versus the instrument response to the analyte. STOCK STANDARD SOLUTION: A concentrated solution containing a single certified standard that is a method analyte, or a concentrated solution of a single analyte prepared in the laboratory with an assay reference compound. Stock standard solutions are used to prepare calibration standards.

SURROGATES: Organic compounds which are similar to analytes of interest in chemical composition, extraction and chromatography, but which are not normally found in environmental samples. These compounds are spiked into all blanks, standards, samples and spiked samples prior to analysis. Percent recoveries are calculated for each surrogate.

TARGET: A software system that combines full processing, reporting and comprehensive review capabilities, regardless of chromatographic vendor and data type.

TARGET DB: An oracle database used to store and organize all Target data files.

KATAHDIN INFORMATION MANAGEMENT SYSTEM (KIMS): A complete multiuser system with the capabilities of integrating laboratory instrumentation, generating laboratory worksheets, providing complete Lab Order status and generating reports. KIMS utilizes these features through a database.

#### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of analysts experienced in the analysis of semivolatile organic compounds by EPA Method 8270. Each analyst must demonstrate and document their ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, "Personnel Training & Documentation of Capability," current revision.

It is the responsibility of all Katahdin technical personnel involved in analysis of semivolatiles by Method 8270 to read and understand this SOP, to adhere to the procedures outlined, and to properly document their data in the appropriate lab notebook. Any deviations from the test or irregularities with the samples should also be recorded in the lab notebook and reported to the Department Manager or designated qualified data reviewer responsible for this data.

It is the responsibility of the Department Manager to oversee that members of their group follow this SOP, to ensure that their work is properly documented and to initiate periodic review of the associated logbooks.

### 1.3 Safety

Users of this procedure must be cognizant of inherent laboratory hazards, proper disposal procedures for contaminated materials and appropriate segregation of

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hazardous wastes. The toxicity or carcinogenicity of each reagent used in this method has not been precisely defined; however, each chemical should be treated as a potential health hazard. A reference file of material safety data sheets is available to all personnel involved in the chemical analysis. Everyone involved with the procedure must be familiar with the MSDSs for all the materials used in this procedure.

Each qualified analyst or technician must be familiar with the Katahdin Analytical Environmental Health and Safety Manual including the Katahdin Hazardous Waste Management Plan and follow appropriate procedures. These include the use of appropriate personal protective equipment (PPE) such as safety glasses, gloves, and lab coats when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location of all safety equipment. Each analyst shall receive a safety orientation from their Department Manager, or designee, appropriate for the job functions they will perform.

### 1.4 Pollution Prevention/Waste Disposal

Whenever possible, laboratory personnel should use pollution prevention techniques to address their waste generation. Refer to the current revision of the Katahdin Hazardous Waste Management Program for further details on pollution prevention techniques.

After analysis, autosampler vials containing sample extracts in methylene chloride are returned to the SVOA hood, and the contents transferred to a labeled waste container. The contents of this container are disposed of in accordance with the Katahdin Hazardous Waste Management Plan and Safety Manual and SOP SD-903, "Sample Disposal," current revision. Expired standards are lab packed, placed in the Katahdin hazardous waste storage area, and disposed of in accordance with this SOP.

#### 2.0 SUMMARY OF METHOD

The process involves the extraction of semivolatiles from a sample using an appropriate solvent followed by clean up steps (where applicable) and concentration of the extract (refer to Katahdin SOP CA-502, "Preparation of Aqueous Samples for Extractable Semivolatile Analyses", SOP CA-512, "Preparation of Sediment/Soil Samples by Sonication Using Method 3550 for Subsequent Extractable Semi-Volatiles Analysis" and SOP CA-526, "Preparation of Sediment/Soil Samples by Soxhlet Extraction Using Method 3540 for Subsequent Extractable Semivolatile Analysis"). An aliquot of the final extract is injected into the gas chromatograph for compound separation by capillary column, followed by the electron impact mass spectrometer for identification and quantitation.

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#### 3.0 INTERFERENCES

Interfering contamination may occur when a sample containing low concentrations of SVOCs is analyzed immediately after a sample containing high concentrations of SVOCs. Any samples that have suspected carryover must be reanalyzed.

#### 4.0 APPARATUS AND MATERIALS

- 4.1 GC: Hewlett Packard 5890 and/or 6890
- 4.2 Mass Spectrometers (MS): HP5975, HP5973, HP5972 and/or HP5970
- 4.3 Helium: Carrier gas for routine applications. All carrier gas lines must be constructed from stainless steel or copper tubing; non-polytetrafluoroethylene (non-PTFE) thread sealant or flow controllers with rubber component are not to be used.
- 4.4 Autosamplers: HP 7673As
- 4.5 Hamilton syringes: 2.00 uL to 10 mL
- 4.6 Volumetric glassware: Grade A or equivalent
- 4.7 Columns: DB-5MS 30m, 0.25mm I.D., 25um film thickness, columns (J&W Scientific) or equivalent.
- 4.8 Acquisition System: The acquisition system must be interfaced to the MS and allow continuous acquisition of data throughout the duration of the chromatographic program. It must permit, at a minimum, the output of time vs. intensity (peak height or peak area). Hewlett Packard Chemstation or equivalent.
- 4.9 Data System: The Target software is used for processing data and generating forms.

### 5.0 REAGENTS

- 5.1 J.T. Baker Ultra Resi-Analyzed methylene chloride (or equivalent)
- 5.2 Purge and trap grade methanol
- 5.3 Standards: Stock standards and working standards are received and recorded in accordance with SOP CA-106 "Standard Preparation and Documentation".

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5.3.1 The expiration date for all standards is one year from date of opening the ampule. If the manufacturer's expiration date is before this one year date, the manufacturer's expiration must be followed. New standards must be opened if degradation is observed.

### 5.3.2 Secondary dilution standards

- 5.3.2.1 The standards are prepared on an as needed basis (or every 6 months) and stored in screw-cap amber bottles with Teflon liners in the BNA standards freezer between uses. Standards prepared from various stock solutions must always use the first expiration date of any of the solutions used for preparation.
- 5.3.2.2 Calibration Mix A Prepare standards in methylene chloride containing the compounds listed in Table 3. The final concentration of each compound is 20 ug/mL.
- 5.3.2.3 Calibration Mix B Some compounds must be calibrated at higher concentrations. For these compounds a secondary standard is prepared which will "boost" the concentration of these compounds in the initial calibration. The concentration of this standard is determined on a project to project basis.
- 5.3.2.4 Internal Standard Solution Prepare standard in methylene chloride containing 1,4-dichlorobenzene-d4, naphthalene-d8, acenaphthene-d10, phenanthrene-d10, chrysene-d12, and perylene-d12 at a final concentration of 80 ug/mL.
- 5.3.2.5 DFTPP Solution Prepare standard in methylene chloride containing DFTPP at a final concentration of 25 ug/mL.
- 5.3.2.6 Independent Calibration Verification (ICV) Standard From a source independent of the calibration standards, prepare a standard in methylene chloride containing the compounds listed in Table 3. The final concentration of each compound is 2 ug/mL.

#### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

All semivolatile sample extracts must be analyzed within forty days following the date of extraction.

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#### 7.0 PROCEDURES

- 7.1 NAMING AND CODING CONVENTIONS FOR ANALYTICAL STANDARDS Used in accordance with SOP CA-106 "Standard Preparation and Documentation".
- 7.2 COMPUTER (DATA SYSTEM) CONVENTIONS -

Conventions for all instruments are as follows:

Sub-Directory for data acquisition and storage: C:\HPCHEM\1\DATA Tune file: DFTPP.U

Method files: LSPSIMXX.M (all samples and standards)

Where:

XX = the calibration number in chronological order

L = instrument ID (Each instrument is given a unique identifier)

DFTPP390.M (DFTPP tuning acquisition)

NOTE: All acquisition parameters must be identical for LSPSIMXX.M and DFTPP2. M.

Data Files: L\_\_\_\_.D, where \_\_\_\_ is a number in chronological order from 0001 to 9999 and L is the instrument ID (Each instrument is given a unique identifier). This file also contains the Quantitation output file.

Data Files for DFTPP: LD\_\_\_.D, where \_\_\_ is a number in chronological order from 001 to 999 and L is the instrument ID (Each instrument is given a unique identifier).

### 7.3 INSTRUMENT SPECIFIC PROCEDURES

It is the policy of the GC/MS group that all data be acquired in the batch mode. The following items must be checked prior to data acquisition in the batch mode:

- Ensure that the proper sequence and tune files are being used.
- Check the autosampler syringe (Is it clean? Does the plunger move freely? etc.), its alignment and make sure the solvent rinse vial is full. Ensure that the knurled nut holding the top of the syringe plunger is tight.
- Look at the batch to be analyzed and check the following:

Make sure that the data files are in numerical order with no duplication and that the method file is the same as that used for ICAL or Continuing Calibration analysis.

Bottle numbers match with the numbers on the autosampler tray.

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After the batch has been deemed free of errors, start the batch by using the "Position and run" command under the SEQUENCE menu in MSTop.

7.4 INSTRUMENT TUNING - Prior to the analysis of any calibration standards, blanks or samples, the GC/MS system must be shown to meet the mass spectral key ion and ion abundance criteria for decafluorotriphenylphosphine (DFTPP) tabulated below. Pentachlorophenol, benzidine and DDT are also present in this standard.

DFTPP Key lons and Ion Abundance Criteria		
Mass	Criteria	
51	30.0-60.0 percent of mass 198	
68	less than 2.0 percent of mass 69	
69	present	
70	less than 2.0 percent of mass 69	
127	40.0 – 60.0 percent of mass 198	
197	less than 1.0 percent of mass 198	
198	base peak, 100 percent of mass 198	
199	5.0-9.0 percent of mass 198	
275	10.0-30.0 percent of mass 198	
365	greater than 1.00 percent of mass 198	
441	present, but less than mass 443	
442	greater than 40.0 percent of mass 198	
443	17.0-23.0 percent of mass 442	

All ion abundances must be normalized to m/z 198, the nominal base peak.

The following are the GC/MS operating conditions for injection of DFTPP.

GC/MS Operating Conditions - D	FTPP
Initial column temperature hold	140°C for 3 minutes
Column temperature program	140-275°C at 15 degrees/minute
Final column temperature hold	275°C
Injection port temperature	280°C
Transfer line/source temperature	285°C
Injector - splitless, valve time	0.18 minutes
EPC	inlet B
Constant flow	ON
Constant flow pressure	10psi
Constant flow temperature	30°C
Vacuum comp.	ON
Run time	10-12 minutes
Scan start time	5.0 minutes
Sample volume	2.0 uL of 25 ng/uL DFTPP solution
Carrier gas	helium at @ 1.0 mL/minute
Mass range	35 to 500 amu
Number of A/D samples	4
GC Peak threshold	500 counts
Threshold	10 counts

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Set up the run on the Enviroquant system using "Edit Sample Log Table". For a more detailed explanation of the Enviroquant software, consult the appropriate manual, Organic Department Manager, or senior chemist within the GC/MS group.

The DFTPP solution must be analyzed once at the beginning of each twelve hour period during which standards and/or samples are analyzed. The 12 hour time period for GC/MS system begins at the moment of injection of the DFTPP analysis. The time period ends after twelve hours has elapsed according to the system clock. The last injection must be accomplished prior to the expiration of 12 hours; conceivably, the run-time of an injection could end after the twelve hours.

When the DFTPP has concluded, the run must be evaluated to determine if sample analysis can proceed. The chromatography and the ion ratios must be examined. The DFTPP run is processed using the current algorithms in the Target software.

The DFTPP tuning standard should also be used to assess the column performance and injection port inertness. Calculate the degradation of DDT to DDE and DDD; it should not exceed 20%. Benzidine and pentachlorophenol should be present at their normal responses, with no evidence of peak tailing. For clients requiring DOD criteria, the tailing factors for these two compounds should not exceed 2.

In order to document the performance of benzidine, pentachlorophenol and DDT, the following procedure must be followed. At the PC, which operates the instrument, load the method TUNETAIL.M into the ENVDA screen. Go into the quant drop down menu and select *calculate/generate report*. When that finishes, select *Qedit quant result*. Each compound can now be evaluated. Double click on benzidine and select *ChromEval* and then *Evaluate tailing*. Follow the instructions given on the screen to evaluate tailing. Send the report to the printer. Repeat the procedure for pentachlorophenol. Repeat the procedure for DDT, selecting *Evaluate degradation*. Follow the instructions given on the screen and then send the report to the printer. The report should be filed with the tune raw data.

If the results indicate the system does not meet acceptance criteria, the GC/MS must be manually tuned. Once the manual tune procedure is completed, DFTPP must be re-injected and reevaluated. If the instrument still does not meet criteria, notify your Department Manager. Under no circumstances should calibration proceed if the instrument DFTPP is not in criteria.

#### 7.5 INSTRUMENT CALIBRATION

#### 7.5.1 Initial Calibration for Method 8270-SIM

Prior to the analysis of samples and required method blanks, and after the instrument DFTPP tuning criteria have been met, the GC/MS system must be calibrated at six different concentrations, typically, 0.20, 0.50, 1.0, 2.0, 5.0 and 8.0 ng/uL. This is done to determine instrument sensitivity and the

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linearity of GC/MS response for the semivolatile target and surrogate compounds.

Some SIM compounds may need to be calibrated at higher concentrations. A second standard is prepared containing these compounds. The two standards are combined as in the example below. The full aliquot is used and spiked with the appropriate amount of IS.

#### Example -

For a calibration at the following levels:

Calibration mix A would be prepared containing ALL analytes at 20 ng/ul Calibration Mix B would be prepared containing phenols and phthalates at 20 ng/ul.

Final PAH conc. (ng/uL)	Final Conc. Phenols and phthalates (ng/ul)	Cal-Mix A Added (uL)	Cal-Mix B Added (uL)	MeCl <sub>2</sub> Added (uL)	Final Volume (uL)
0.20	1.0	10	40	950	1000
0.50	2.0	25	75	900	1000
1.0	3.0	50	100	850	1000
2.0	4.0	100	100	800	1000
5.0	5.0	250	0	750	1000
8.0	8.0	400	0	600	1000

Note: Calibration Mix B only is used to boost the phenols and phthalates concentrations in Cal. levels 1 through 4.

The GC/MS operating conditions for the calibration standards injections are the same as for the DFTPP with the following exceptions:

GC/MS Operating Conditions – Calibration and Samples				
Column temperature program	40°C for 3 min. to 300°C at 10°/min.			
Final column temperature hold	300°C			
	35 minutes (time may vary dependent			
Run time	upon column length)			
	2.0-6.0 minutes (time may vary			
	dependent			
Scan start time	upon column length)			
Sample volume	2 uL			

The conditions are set up in the method file LSPSIMXX.M

After analysis of the six calibration points, they must be quantitated and evaluated for adherence to QC criteria. Minimum requirements of ID files are the use of specific quantitation ions and quantitating a specific set of targets and surrogates with a set internal standard. Of particular importance when

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performing SIM analysis are the ion ratios. These requirements are found in Tables 3 and 5.

#### 7.5.2 Initial Calibration Criteria

Relative response factors (RRFs) must be calculated and evaluated for each target compound and surrogate. The RRF is defined as follows:

$$RRF = \underbrace{Ax}_{A_{IS}} X \underbrace{C_{IS}}_{Cx}$$

area of the primary ion for the target compound where: Ax =

area of the primary ion for the corresponding istd

 $A_{IS} = C_{IS} = 0$ concentration of the istd (ng/uL) concentration of the target compound

After the calibration points have been quantitated, update the calibration curve points using the Target data processing software to generate the RRF's and %RSD's for all analytes. If information is needed concerning the use of these programs, consult the Organic Department Manager or a senior chemist within the group.

Response factor criteria have been established for the calibration of the semivolatile target and surrogate compounds. These criteria must be met in order for the calibration curve to be considered valid. The percent RSD for each calibration check compound (CCC) must be less than or equal to 30 percent. There are three CCC's: Acenaphthene, Fluoranthene, and Benzo(a)pyrene. There are no criteria for the SPCC compounds. This is also applicable to clients that request DOD criteria.

7.5.2.1 Linearity of Target Analytes (This is also applicable to clients that request DOD criteria.)

If the RSD of any target analyte is 15% or less, then the response factor is presumed to be constant over the calibration range, and the average response factor may be used for quantitation.

If the RSD of any target analyte exceeds 15%, then a calibration option outlined in section 7.0 of method 8000 will need to be employed.

Option 1 (Section 7.5.2 of method 8000 - Rev. 2, 12/96), is a linear regression of instrument response versus the standard concentration. The correlation coefficient (r) for each target analyte and surrogate must be greater than or equal to 0.995. Target software calculates the correlation coefficient squared (r2). This must be equal to or greater than 0.990.

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Option 2 (Section 7.5.3 of method 8000 - Rev. 2, 12/96), is a non-linear calibration model not to exceed a third order polynomial. The lab would use a quadratic model or second order polynomial. The use of a quadratic model requires six calibration points. In order for the quadratic model to be acceptable, the coefficient of determination must be greater than or equal to 0.990.

If time remains in the clock after meeting the initial calibration acceptance criteria, samples may be analyzed. The calibration must be verified each twelve hour time period (time period starts from the moment of the DFTPP injection) for Method 8270-SIM. The SSTD1.0 in the curve may be used as the continuing calibration standard as long as it meets the continuing calibration acceptance criteria. All sample results must be quantitated using the initial calibration response factors.

7.5.2.2 Immediately following calibration an Independent Calibration Verification Standard must be analyzed. For clients requiring DOD criteria, all project analytes must be within +/- 20% of true value.

#### 7.5.3 Continuing Calibration

A check of the calibration curve must be performed once every twelve hours immediately following analysis of the tuning compound DFTPP. This check contains all target compounds and surrogates at a concentration of 1.0 ng/uL.

After quantitation of the 1.0 ng/uL continuing calibration check, response factors must be calculated and compared to the average response factors in the initial calibration. The Target program calculates the calibration check response factors and compares them to the average RFs in the calibration curve by calculating percent differences. The method 8270 CCC's must have a % difference of +/- 20%D in order to be considered in criteria. These conditions must be met before method blank and/or sample analysis can begin. For clients requiring DOD criteria, all project analytes and surrogates must be within +/- 20%.

If the continuing calibration check does not meet criteria, corrective action must be taken. Depending on the situation, corrective action may be as follows:

- Re-analyze the 1.0 ng/uL continuing calibration check.
- Change the septum; clean the injection port; install a clean, silanized quartz liner; cut off a small portion (1" to 3") of the front end of the capillary column. This is usually performed when chromatography is poor. Record any of these actions in the appropriate instrument maintenance logbook.
- Analyze a new initial calibration curve.

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The last option, the generation of a new initial calibration curve, is usually chosen when percent difference are >30%. In these instances, there is little or no chance of a continuing calibration reanalysis meeting criteria. If there is any doubt concerning which corrective action to undertake, consult the Organic Department Manager or a senior chemist within the group.

If the continuing calibration does meet the criteria specified above then analysis may precede using initial calibration response factors.

#### 7.5.4. Retention Time Windows

Retention time windows for each analyte and surrogate are set at the midpoint standard of the calibration curve, following every ICAL. On days when an ICAL is not performed, the initial CCV may be used. For each sample, the RRT shall be compared with the mid-point of the ICAL or the most recently updated RRT. If the RRT has changed by more than  $\pm$  0.006 RRT units indicates a significant change in system performance and the laboratory must take appropriate corrective action.

For projects or clients requiring DoD QSM 4.1, IS EICP areas must be within -50% to + 100% of the ICAL midpoint standard. The retention time must be  $\pm$  30 seconds from the retention time of the ICAL midpoint standard.

#### 7.6 SAMPLE ANALYSIS

Sample extracts may be analyzed only after the GC/MS system has met tuning criteria, initial calibration and continuing calibration requirements. Ensure that the same instrument conditions are being used for tuning, calibration and sample analysis by reviewing the GC parameters using the "Edit entire method" option under the Method menu in MSTOP. Note that you can not edit a method if the instrument is running.

Extracts are stored in the refrigerator in the organics extraction laboratory at 4°C ±2°C. Remove them from the refrigerator and place them in the GC/MS laboratory semivolatile hood when ready for analysis.

Prepare a 1.8 mL clear glass vial (crimp top) with a disposable insert (350 uL). Add 100 uL of sample extract and 1.0 uL of the 80 ng/uL IS stock to the vial and then cap. This gives a 0.8 ng/uL final concentration for the internal standard compounds. The samples are topped with Teflon lined crimp top caps.

#### 7.7 FINAL DATA PACKAGE

#### 7.7.1 Initial Data Review (IDR)

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The initial data review is accomplished by the analyst who ran the samples and is a review of sufficient quality and detail to provide a list of samples that need to be reanalyzed or diluted and reanalyzed. The initial data review is performed on the detailed quantitation reports of the analyzed sample. This data review examines criteria that directly impact whether or not the sample needs to be reanalyzed:

- Surrogate Recoveries
- Internal Standard Area Stability
- Method Blank Acceptance
- Chromatography
- Target Compound Detection/Quantitation/Review for false positives
- Laboratory Control Sample Recoveries
- Matrix Spike/Matrix Spike Duplicate Recoveries

The analyst must evaluate all data using the QA Acceptance Criteria table found within this SOP (Table 1). This table gives acceptance criteria and corrective actions for criteria that are not met. In addition to evaluating QC elements, the chromatography and quantitation of target analytes must be reviewed. During this review, the analyst checks the integration of each individual peak. The hardcopy has false positives crossed out so they can be reviewed for appropriateness by the Organic Department Manager.

### 7.7.2 Chromatography

The chromatography should be examined for the presence or absence of any ghost peaks and can also be used as an indication of whether or not matrix interferences might be affecting surrogate recoveries and/or istd area recoveries. Whether or not the chromatography is acceptable is a judgment call on the part of the analyst and should be used in conjunction with other monitored QC (e.g. surrogate recoveries) to determine the necessity of reanalyzing.

Manual integrations are to be performed when chromatographic conditions preclude the computer algorithm from correctly integrating the peak of concern. In no instance shall a manual integration be performed solely to bring a peak within criteria.

Each peak of concern is examined by the primary analyst to ensure that the peak was integrated properly by the computer algorithm. Should a manual integration be necessary (for instance, due to a split peak, peak tailing, or incomplete resolution of isomeric pairs), an "m" qualifier will automatically be printed on the quantitation report summary.

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This manual integration package must then be submitted to the Department Manager or his/her designee, who will review each manual integration.

For specific manual integration procedures, refer to Katahdin SOP QA-812, "Manual Integration", current revision.

### 7.7.3 Target Compound Detection/Quantitation

The semivolatile ID files have been set up to err on the side of false positives; that is to identify and quantitate peaks as target compounds that may not necessarily be valid hits. It is the responsibility of the GC/MS analyst to use his/her technical judgment to determine if the identification of a target compound is correct or not.

If any target concentration exceeds the upper limit, a dilution must be made and analyzed. The dilution chosen should keep the concentration of the largest target compound hit in the upper half of the initial calibration range. LCS and MS/MSD samples need not be diluted to get spiked analytes within the calibration range.

The requirements for qualitative verification by comparison of mass spectra are as follows:

- All ions present in the standard mass spectra at a relative intensity > 10% must be present in the sample spectrum.
- The relative intensities of primary and secondary ions must agree within ±20% between the standard and sample spectra.
- lons greater than 10% in the sample spectrum but not present in the standard spectrum must be considered and accounted for by the analyst.

If a compound cannot be verified by all three criteria above, but, in the technical judgment of the mass spectral interpretation specialist, the identification is correct, then the laboratory shall report that compound on the Form 1 as a valid hit.

The GC/MS laboratory initial data review must be completed within twelve hours of batch completion; in the majority of instances, the initial review should be accomplished at the beginning of a work shift for the previous set of analyses.

#### 7.7.4 Reporting

After the chromatograms have been reviewed and any target analytes have been quantitated using Target, the necessary files are brought into QuickForms. Depending on the QC label requested by the client, a Report

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of Analysis (ROA) and additional reports, such as LCS forms and chronology forms, are generated. The package is assembled to include the necessary forms and raw data. The data package is reviewed by the primary analyst and then forwarded to the secondary reviewer. The secondary reviewer validates the data and checks the package for any errors. When completed, the package is sent to the department manager for final review. A complete review checklist is provided with each package. The final data package from the Organics department is then processed by the Data Management department.

### 7.8 Injection Port Liner Cleaning And Silanizing Procedure

- Remove the rubber o-ring from the liner and place the liner in a large Erlenmeyer flask.
- In the hood, pour nitric acid into the flask until the liner is covered. Place the flask on a hotplate and boil for 2-3 hours.
- Let cool; drain nitric acid and thoroughly flush the liner with water.
- Bake briefly in the muffle oven until liner is dry and cool to room temperature.
- Place the liner in a beaker, fill with Sylon and let it soak for at least two hours.
- Take out the liner and rinse it thoroughly with toluene.
- Rinse the liner thoroughly with purge and trap grade methanol.
- Bake the liner in the muffle oven for a minimum of three hours.

#### 7.9 Instrument Maintenance

Instrument preventative maintenance is performed on a semi-annual basis by GC/MS chemists. This maintenance includes a thorough inspection and cleaning of all parts, including changing rough and turbopump oils. GC/MS analysts perform other maintenance on an as-needed basis. Typically, routine maintenance involves clipping off the front end of the DB-5MS column, replacing the injection port septum, and installing a freshly silanized quartz liner after sample analysis.

All maintenance must be documented in the instrument-specific maintenance log, whether it is routine or not. The Department Manager must authorize any maintenance over and above a routine source cleaning.

#### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

Refer to Table 1 and to details in this section for a summary of QC requirements, acceptance criteria, and corrective actions. These criteria are intended to be guidelines for analysts. The criteria does not cover all possible situations. If any of the QC requirements are outside the recovery ranges listed in this section or in Table 1, all associated samples must be evaluated against all the QC. In some cases data may be reported, but may be reanalyzed in other cases. Making new reagents and standards may be necessary if the standardization is suspect. The corrective actions listed in this section and in Table 1 may rely on analyst experience to make sound scientific judgments. These decisions are

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based on holding time considerations, client and project specific Data Quality Objectives and on review of chromatograms. The supervisor, Operations Manager, and/or Quality Assurance Officer may be consulted to evaluate data. Some samples may not be able to be reanalyzed within hold time. In these cases "qualified" data with narration may be advisable after consultation with the client.

In some cases the standard QC requirements listed in this section and in Table 1 may not be sufficient to meet the Data Quality Objectives of the specific project. Much of the work performed at the lab is analyzed in accordance with specific QC requirements spelled out in a project specific Quality Assurance Project Plan (QAPP) or in a program specific Quality Systems Manual (QSM). The reporting limits, acceptance criteria and/or corrective actions may be different than those specified in this SOP. In these cases the appropriate information will be communicated to the Department Manager and/or senior chemists before initiation of the analyses so that specific product codes can be produced for the project. In addition, the work order notes for each project will describe the specific QAPP or QSM to be followed.

#### 8.1 Method Blank Criteria

A method blank is defined as a volume of a clean reference material (deionized distilled water for water samples, baked organic-free sand for soil/sediment matrices) that is carried through the entire analytical procedure. One method blank must be extracted with each group of samples of a similar matrix and must be analyzed on the GC/MS system that was used to analyze the samples.

An acceptable method blank must contain less than or equal to the PQL of any target compound. For clients requiring DOD criteria, no analytes detected at  $> \frac{1}{2}$  PQL and  $> \frac{1}{10}$  the amount measured in any sample or  $\frac{1}{10}$  the regulatory limit.

If the method blank exceeds these contamination levels, the analytical system is considered out of control and corrective action must be taken before sample analysis.

Reanalysis of the blank is the first step of the corrective action; if that does not solve the problem, a Katahdin Corrective Action Report (CAR) will be initiated. Corrective action will be specified after consultation including the Department Manager, Operations Manager, and QA Officer.

#### 8.2 Surrogate Recoveries

The five surrogates (2-Methylnaphthalene-d10, 2,4-Dibromophenol, Fluorene-d10, Pyrene-d10 and 1,4-Dioxaned8) must meet the current statistically derived or nominal acceptance limits. If statistical limits have not been established then the surrogate recovery must meet the nominal limits of 30-150%. For clients requiring DOD criteria, use acceptance limits specified by DOD or use in-house limits where none are specified.

If specifications are not met, the sample (or blank) should be reanalyzed. If specifications are met in the reanalysis, this reanalysis should only be submitted. If

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surrogate specifications are not met in the sample or method blank reanalysis, a Corrective Action Report (CAR) should be initiated. Corrective action will be specified after consultation including the Department Manager and Operations Manager.

For further information regarding the acceptance of surrogate recoveries, consult the Organic Department Manager.

### 8.3 Internal Standard Responses

Internal standard responses and retention times (RT) in all samples and blanks must be evaluated as part of the technical data review. The method files have been set up to only detect compounds that fall within a set RT window. For Method 8270-SIM analysis, if the extracted ion current profile (EICP) area for any internal standard changes by more than a factor of two (-50% to +100%) as compared to the daily continuing calibration standard, reanalysis must occur. If the reanalysis meets criteria, only the in-criteria run should be reported. If the reanalysis is still out of criteria, both analyses should be included in the sample package set.

For projects or clients requiring DoD QSM 4.1, IS EICP areas must be within -50% to  $\pm$  100% of the ICAL midpoint standard. The retention time must be  $\pm$  30 seconds from the retention time of the ICAL midpoint standard.

MS/MSD samples that do not meet the EICP area criteria above do not have to be reanalyzed.

#### 8.4 Laboratory Control Sample (LCS)

An LCS must be performed for each group of samples of a similar matrix, for the following, whichever is more frequent:

- Every 20 samples of a similar matrix or similar concentration, or
- Every batch of samples extracted.

Statistical limits are compiled annually for LCS recoveries (archived in QA office). Statistical limits are only calculated when at least 20 usable data points are obtained for any given compound. If insufficient data points are available, nominal limits are set by the section supervisor, Laboratory Operations Manager and Quality Assurance Officer. Refer to Katahdin SOP QA-808, "Generation and Implementation of Statistical QC Limits and/or Control Charts", current revision.

The use of statistical limits versus nominal limits is dependent on the client and project. This information is communicated to the Organic Department Manager through the Katahdin project manager. It is standard practice to use statistical limits for reporting purposes and to evaluate any QC criteria exceedances. However, nominal limits of 30-150% may be used for some projects or states (i.e. South Carolina). For clients requiring DOD criteria, use acceptance limits specified by DOD or use in-house limits where none are specified.

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The LCS recoveries for all analytes are evaluated. All of the compounds of interest must fall within the established statistical limits or nominal limits with the following sporadic exceedance allowances, for DoD clients.

# of Analytes	# of Allowable Exceedances
> 90	5
71 – 90	4
51 – 70	3
31 – 50	2
11 – 30	1
<11	0

If less than the number of allowable exceedances fail the statistical limits, no corrective action is needed. If greater than the number of allowable exceedances fail the statistical limits, corrective action may be taken. Corrective actions may vary with each situation. However, in the case where the failures are high and the samples are non-detect for those compounds, then no corrective action is required. Otherwise, corrective action may involve reanalysis or recalibration. The specific corrective actions taken will rely on analyst experience to make sound scientific judgments while considering client objectives, other quality control indicators and/or the ability to reanalyze a sample within holding time.

For non-DoD clients corrective action is only taken if greater than 10% of the analytes of interest are outside of the laboratory established acceptance limits.

#### 8.5 Matrix Spike/Matrix Spike Duplicate (MS/MSD) Criteria

Matrix Spike and Matrix Spike Duplicates must be extracted and analyzed for each group of up to 20 samples of a similar matrix or similar concentration. In the event insufficient sample volume is available an LCS/LCS Duplicate is extracted and analyzed in place of the MS/MSD.

Statistical limits are compiled annually for MS/MSD recoveries for a short list of the spiked compounds (Acenaphthene, Pentachlorophenol and Pyrene). Nominal limits of 30-130% are used for all other compounds. Generally, corrective action is only taken for the short list of the spiked compounds. The specific corrective actions will rely on analyst experience to make sound scientific judgments while considering client objectives, other quality control indicators and/or the ability to reanalyze a sample within holding time. For clients requiring DOD criteria, use acceptance limits specified by DOD or use in-house limits where none are specified.

A Corrective Action Report (CAR) must be filled out and filed if any criteria for percent recovery or relative percent difference are not met to document any decisions with reporting data.

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#### 9.0 METHOD PERFORMANCE

The method detection limit (MDL) is defined as the minimum concentration of a substance that can be measured and reported with 99% confidence that the value is above zero. The MDLs shall be determined and verified one time per type of instrument unless otherwise required by the method.

A Limit of Detection (LOD) is an estimate of the minimum amount of a substance that an analytical process can reliably detect. An LOD is analyte and matrix specific and may be laboratory-dependent. LODs must be determined for all parameters for which the laboratory is accredited under the DoD Environmental Laboratory Accreditation Program. LOD's must be verified for every preparation and analytical method combination and on every applicable instrument on a quarterly basis.

The Limit of Quantitation (LOQ) is the minimum levels, concentrations, or quantities of a target variable (e.g., target analyte) that can be reported with a specified degree of confidence. The LOQ shall be set at the lowest point in the calibration curve for all analyses utilizing an initial calibration. LOQ's must be verified quarterly for every preparation and analytical method combination and on every applicable instrument on a quarterly basis for all parameters included in the DoD Scope of Accreditation. The LOQ must be verified at least once annually if the analysis is not included in the DoD Scope of Accreditation.

MDLs are filed with the Organic Department Manager and then with the QAO. LOD and LOQ verifications are filed with the QAO

Refer to the current revision of Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, for procedures on determining the MDL.

Refer to the current revision of Method 8270 for other method performance parameters and requirements.

#### 10.0 APPLICABLE DOCUMENTS/REFERENCES

"Test Methods for Evaluating Solid Wastes: Physical/Chemical Methods." SW-846, 3rd edition, Final Updates I, II, IIA, IIB, III, IIIA, and IIIB, Nov 2004, Method 8270C.

Test Methods for Evaluating Solid Waste, Physical/Chemical Methods, USEPA SW846, 3<sup>rd</sup> Edition, Final Updates I, II, IIA, IIB, III, IIIA, IIB and IV, February 2007, Method 8270D.

Department of Defense Quality Systems Manual for Environmental Laboratories (DoD QSM), Current Version.

The National Environmental Laboratory Accreditation Conference (NELAC) Standards, June 2003.

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The NELAC Institute, Laboratory Accreditation Standards, Volume 1, Management and Technical Requirements for Laboratories Performing Environmental Analysis, 10/06/2010.

Katahdin SOP CA-101, Equipment Maintenance and Troubleshooting, current revision.

Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, current revision.

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## TABLE 1

# QC REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action
Check of mass spectral ion intensities using DFTPP	Prior to initial calibration and calibration verification	Refer to the criteria listed in Section 7.4	Retune instrument, and verify
Six-point initial calibration for all analytes	Initial calibration prior to sample analysis	RSD ≤30 for RFs of the CCCs; Average %RSD < 15% for all compounds. Refer to section 7.5.2.1 for more details.	Repeat calibration if criterion is not met
Independent calibration verification	Once after Initial calibration	± 20 % D	Reanalyze standard     Reprep standard     Reprep standard from fresh stock.
Continuing calibration verification	Once per each 12 hours, prior to sample analysis	CCCs ≤ 20%D	Repeat initial calibration and reanalyze all samples analyzed since the last successful calibration verification
ISs	Immediately after or during data acquisition of calibration check standard	Retention time ± 30 seconds; EICP area within -50% to +100% of last calibration verification (12 hours) for each IS	Inspect mass spectrometer or GC for malfunctions; mandatory reanalysis of samples analyzed while system was malfunctioning
Demonstration of ability to generate acceptable accuracy and precision	Once per analyst initially and annually thereafter	All recoveries within method QC acceptance limits.	Recalculate results; locate and fix problem; reextract/reanalyze P&A study for those analytes that did not meet criteria
Method blank	One per prep batch	No analytes detected > PQL	(1) Investigate source of contamination (2) Evaluate the samples and associated QC: i.e.If the blank results are above the PQL, report samples that are <pql or=""> 10X the blank result. Reprep a blank and the remaining samples.</pql>
LCS for all analytes	One LCS per prep batch	Statistically derived from lab data or nominal limits depending on the project See also section 8.4 of this SOP for more information on allowable exceedances.	(1) Evaluate the samples and associated QC: i.e. If an MS/MSD was performed and acceptable, narrate. If an LCS/LCSD was performed and only one was unacceptable, narrate. If the surrogate recoveries in the LCS are low but are acceptable in the blank and samples, narrate. If the LCS rec. is high but the sample results are <pql, a="" and="" blank="" narrate.="" otherwise,="" remaining="" reprep="" samples.<="" td="" the=""></pql,>

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# TABLE 1 (cont.)

# QC REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action
Surrogate spike	Every sample, control, standard, and method blank	Statistically derived limits.	(1) Check chromatogram for interference; if found, flag data (2) If not found, check instrument performance; if problem is found, correct and reanalyze (3) If still out reextract and analyze sample (4) If reanalysis is out, flag data
MS/MSD	One MS/MSD per every 20 samples	Statistically derived from lab data or nominal limits depending on the project. Nominal limits are used as default limits.	(1) Evaluate the samples and associated QC: i.e. If the LCS results are acceptable, narrate. (2) If both the LCS and MS/MSD are unacceptable reprep the samples and QC.
MDL and/or LOD/LOQ Verification study		806, "Method Detection Limit, Inst cations", current revision.	rument Detection Limit and Reporting

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## TABLE 2

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Demonstrate acceptable analytical capability	Prior to using any test method and at any time there is a significant change in instrument type, personnel, test method, or sample matrix.	QC acceptance criteria published by DoD, if available; otherwise, method-specific criteria.	Recalculate results; locate and fix problem, then rerun demonstration for those analytes that did not meet criteria.	NA.	This is a demonstration of analytical ability to generate acceptable precision and bias per the procedure in Appendix C. No analysis shall be allowed by analyst until successful demonstration of capability is complete.
LOD determination and verification LOQ	Refer to current revision of SOP QA-806 Refer to current				
establishment and verification	revision of SOP QA-806				
Tuning	Prior to ICAL and at the beginning of each 12-hour period.	Refer to method for specific ion criteria.	Retune instrument and verify. Rerun affected samples.	Flagging criteria are not appropriate.	Problem must be corrected. No samples may be accepted without a valid tune.
Breakdown check (DDT Method 8270 only)	Correct problem then repeat breakdown check.	Degradation ≤ 20% for DDT. Benzidine and pentachlorophenol should be present at their normal responses, and should not exceed a tailing factor of 2.	At the beginning of each 12-hour period, prior to analysis of samples.	Flagging criteria are not appropriate.	No samples shall be run until degradation ≤ 20%.
Minimum five- point initial calibration (ICAL) for all analytes	ICAL prior to sample analysis.	1. Average response factor (RF) for SPCCs ≥ 0.050.  2. RSD for RFs for CCCs ≤ 30% and one option below: Option 1: RSD for each analyte ≤ 15%; Option 2: linear least squares regression r ≥ 0.995; Option 3: non-linear regression—coefficient of determination (COD) r2 ≥ 0.99 (6 points shall be used for second order).	Correct problem then repeat ICAL.	Flagging criteria are not appropriate.	Problem must be corrected. No samples may be run until ICAL has passed. Calibration may not be forced through the origin.
Second source calibration verification (ICV)	Once after each ICAL.	All project analytes within ± 20% of true value.	Correct problem and verify second source standard. Rerun second source verification. If that fails, correct problem and repeat ICAL.	Flagging criteria are not appropriate.	Problem must be corrected. No samples may be run until calibration has been verified.

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# TABLE 2 (cont.)

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Retention time window position establishment for each analyte and surrogate	Once per ICAL.	Position shall be set using the midpoint standard of the ICAL curve when ICAL is performed. On days when ICAL is not performed, the initial CCV is used.	NA.	NA.	
Evaluation of relative retention times (RRT)	With each sample.	RRT of each target analyte within ± 0.06 RRT units.	Correct problem, then rerun ICAL.	Flagging criteria are not appropriate.	Laboratories may update the retention times based on the CCV to account for minor performance fluctuations or after routine system maintenance (such as column clipping). With each sample, the RRT shall be compared with the most recently updated RRT. If the RRT has changed by more than ±0.06 RRT units since the last update, this indicates a significant change in system performance and the laboratory must take appropriate corrective actions as required by the method and rerun the ICAL to reestablish the retention times.
Continuing calibration verification (CCV)	Daily before sample analysis and every 12 hours of analysis time.	1. Average RF for SPCCs ≥ 0.050. 2. %Difference/Drift for all target compounds and surrogates ≤ 20%D (Note: D = difference when using RFs or drift when using least squares regression or non-linear calibration).	DoD project level approval must be obtained for each of the failed analytes or corrective action must be taken. Correct problem, then rerun calibration verification. If that fails, then repeat ICAL. Reanalyze all samples since last acceptable CCV.	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply Q-flag to all results for the specific analyte(s) in all samples since last acceptable CCV.	Problem must be corrected. Results may not be reported without a valid CCV. Flagging is only appropriate in cases where the samples cannot be reanalyzed.
Internal standards verification	Every field sample, standard, and QC sample.	Retention time ± 30 seconds from retention time of the midpoint standard in the ICAL; EICP area within -50% to +100% of ICAL midpoint standard.	Inspect mass spectrometer and GC for malfunctions. Reanalysis of samples analyzed while system was malfunctioning is mandatory.	If corrective action fails in field samples, apply Q-flag to analytes associated with the noncompliant IS. Flagging criteria are not appropriate for failed standards.	Sample results are not acceptable without a valid IS verification.

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# TABLE 2 (cont)

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Method blank	One per preparatory batch.	No analytes detected > ½ RL (> RL for common lab contaminants) and > 1/10 the amount measured in any sample or 1/10 the regulatory limit (whichever is greater). Blank result must not otherwise affect sample results.	Correct the problem. Report sample results that are <lod or="">10x the blank concentration. Reprepare and reanalyze the method blank and all associated samples with results &gt; LOD and &lt; 10x the contaminated blank result. Contact Client if samples cannot be reprepped within hold time.</lod>	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply B-flag to all results for the specific analyte(s) in all samples in the associated preparatory batch.	Problem must be corrected. Results may not be reported without a valid method blank. Flagging is only appropriate in cases where the samples cannot be reanalyzed.
LCS containing all analytes to be reported, including surrogates	One per preparatory batch.	The laboratory shall use laboratory control limits (CLs) or use DoD-generated LCS CLs, if available depending on project requirements. Inhouse CLs may not be greater than ± 3 times the standard deviation of the mean LCS recovery. A number of analytes may fall outside the CL but within marginal exceedance limit depending on the total number of analytes in the LCS.	Correct problem, then reprep and reanalyze the LCS and all samples in the associated preparatory batch for failed analytes, if sufficient sample material is available. Refer to Table G-1 for number of marginal exceedences allowed. Contact Client if samples cannot be reprepped within hold time.	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply Q-flag to specific analyte(s) in all samples in the associated preparatory batch.	Problem must be corrected. Results may not be reported without a valid LCS. Flagging is only appropriate in cases where the samples cannot be reanalyzed.
Matrix Spike (MS)	One per preparatory batch per matrix if sufficient sample is available.	The laboratory shall use laboratory LCS CLs or use DoD-generated LCS CLs, if available depending on project requirements.	Examine the project- specific DQOs. Contact the client as to additional measures to be taken.	For the specific analyte(s) in the parent sample, apply J-flag if acceptance criteria are not met.	For matrix evaluation only. If MS results are outside the LCS limits, the data shall be evaluated to determine the source of difference and to determine if there is a matrix effect or analytical error.
Matrix spike duplicate (MSD) or sample duplicate	One per preparatory batch per matrix if sufficient sample is available.	MSD: For matrix evaluation, use laboratory LCS CLs or use DoDgenerated LCS CLs, if available depending on project requirements. MS/MSD: RPD ≤ 30%.	Examine the project- specific DQOs. Contact the client as to additional measures to be taken.	For the specific analyte(s) in the parent sample, apply J-flag if acceptance criteria are not met.	The data shall be evaluated to determine the source of difference.

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# TABLE 2 (cont)

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Surrogate spike	All field and QC samples.	The laboratory shall use laboratory surrogate CLs or use DoD-generated surrogate CLs, if available depending on project requirements.	For QC and field samples, correct problem then reprep and reanalyze all failed samples for failed surrogates in the associated preparatory batch, if sufficient sample material is available. If obvious chromatographic interference with surrogate is present, reanalysis may not be necessary. Contact Client if samples cannot be reprepped within hold time.	Apply Q-flag to all associated analytes if acceptance criteria are not met.	Alternative surrogates are recommended when there is obvious chromatographic interference.
Results reported between DL and LOQ	NA.	NA.	NA.	Apply J-flag to all results between DL and LOQ.	

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# TABLE 3 SUMMARY OF METHOD MODIFICATIONS

TOPIC	KATAHDIN SOP CA-213-11	METHOD 8270, current revision
Apparatus/Materials	none	
Reagents	none	
Sample preservation/ handling	none	
Procedures	none	
QC - Spikes	none	
QC - LCS	none	
QC - Accuracy/Precision	none	
QC - MDL	none	

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# TABLE 4 ANALYTE QUANITIATION AND INTERNAL STANDARDS

Internal Standard: 1,4-dichlorobenzene-d4	2,4-Dinitrotoluene
Target and Surrogates:	2,4-Dinitrophenol
1,4-Dioxane	2,3,4,6-Tetrachlorophenol
1,4-Dioxane-d8 (surrogate)	Diethylphthalate
Benzaldehyde	4-Chlorophenyl-phenyl ether
Phenol	4,6-Dinitro-2-methylphenol
bis(2-Chloroethyl)ether	N-nitrosodiphenylamine
2-Chlorophenol	2-Nitroaniline
2-Methylphenol	3-Nitroaniline
3&4-Methylphenol	4-Nitroaniline
2,2'-Oxybis(1-chloropropane)	Dibenzofuran
Nitrobenzene	4-Nitrophenol
Hexachloroethane	Internal Standard: Phenanthrene-d10
Acetophenone	Target and Surrogates:
N-nitroso-di-n-propylamine	Pentachlorophenol
Internal Standard: Naphthalene-d8	1-Methylphenanthrene (dredge)
Target and Surrogates:	Phenanthrene
Naphthalene	Hexachlorobenzene (special)
1-Methylnaphthalene (dredge)	Anthracene
2-Methylnaphthalene	Fluoranthene
2-Methylnaphthalene-D10 (surrogate)	Carbazole
Isophorone	Di-n-butylphthalate
2-Nitrophenol	4-Bromophenyl-phenyl ether
2,4-Dimethylphenol	Atrazine
bis(2-Chloroethoxy)methane	Internal Standard: Chrysene-d12
2,4-Dichlorophenol	Target and Surrogates:
4-Chloroaniline	Butylbenzylphthalate
Hexachlorobutadiene	3.3'-Dichlorobenzidine
Caprolactam	Pyrene
4-Chloro-3-methylphenol	Benzo(a)Anthracene
Internal Standard: Acenaphthene-d10	Chrysene
Target and Surrogates:	Bis-(2-ethylhexyl)phthalate
1,1'-Biphenyl (dredge)	Pyrene-d10 (surrogate)
2,6 Dimethylnapthalene (dredge)	Internal Standard: Perylene-d12
Acenaphthylene	Target and Surrogates:
Acenaphthene	Perylene (dredge)
Fluorene	Benzo(b)fluoranthene
2-Fluorene-d10 (surrogate)	Benzo(k)fluoranthene
2,4-Dibromophenol (surrogate)	Benzo(e)pyrene (dredge)
2-Chloronaphthalene	Di-n-octylphthalate
Hexachlorocyclopentadiene	Benzo(a)pyrene
2,4,6-Trichlorophenol	Indeno(1,2,3-cd)pyrene
2,4,5-Trichlorophenol	Dibenz(a,h)anthracene
Dimethylphthalate	Benzo(ghi)perylene

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TITLE: ANALYSIS OF SEMIVOLATILE ORGANIC COMPOUNDS BY: SW 846 METHOD 8270 – Modified for Selected Ion Monitoring (SIM)

## TABLE 5

## PROCEDURE CONDENSATION

Clock

12 hours from injection of 50ng DFTPP.

Calibration Curve Criteria

<30% RSD for CCCS <15% RSD average for all analytes in calibration standard

Continuing Calibration Check Criteria

<20% D for CCC compounds

**Additional QC** 

LCS every extraction batch MS/MSD every 20 samples

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# TITLE: ANALYSIS OF SEMIVOLATILE ORGANIC COMPOUNDS BY: SW 846 METHOD 8270 – Modified for Selected Ion Monitoring (SIM)

TABLE 6
SVOA COMPOUNDS AND CHARACERISTIC IONS

COMPOUND	PRIMARY ION	SECONDARY IONS		
1,4-Dioxaned8	96	66		
1,4-Dioxane	88	58		
Benzaldehyde	77	105,106		
Phenol	94	65,66		
bis(2-Chloroethyl)ether	93	63,95		
2-Chlorophenol	128	64,130		
1,4-Dichlorobenzene-d4 (IS)	152	150,115		
2,2'-Oxybis(1-choropropane)	45	77,121		
2-Methylphenol	108	107,77		
Acetophenone	105	77,51		
N-nitroso-di-n-propylamine	70	52,101		
Hexachloroethane	117	201,199		
3&4-Methylphenol	108	107,77		
Nitrobenzene	77	123,51		
Isophorone	82	54,138		
2-Nitrophenol	139	109,81		
2,4-Dimethylphenol	107	122,121		
bis(2-Chloroethoxy)methane	93	63,123		
2,4-Dichlorophenol	162	164,98		
Naphthalene-d8 (IS)	136	137,134		
Naphthalene	128	129,127		
4-Chloroaniline	127	129		
Hexachlorobutadiene	225	223,227		
Caprolactam	113	55,56		
4-Chloro-3-methylphenol	107	77,142		
2,4-Dibromophenol (surr)	252	63,143		
2-Methylnaphthalene-d10 (surr)	152	150		
2-Methylnaphthalene	142	141,115		
1-Methylnaphthalene	142	141,115		
Hexachlorocyclopentadiene	237	235,239		
2,4,6-Trichlorophenol	196	198,200		
2,4,5-Trichlorophenol	196	198,200		
2-Chloronaphthalene	162	127,164		
1,1'-Biphenyl	154	153,76		
2-Nitroaniline	65	92,138		
Dimethylphthalate	163	194,164		
2,6-Dinitrotoluene	165	63,89		
Acenaphthylene	152	151,153		
Acenaphthene	152	154,152		
Acenaphthene-d10 (IS)	164	162		
3-Nitroaniline	138	65,92		
2,4-Dinitrophenol	184	107		
Dibenzofuran	168	139		
DINCHZUIUIAII	100	138		

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# TITLE: ANALYSIS OF SEMIVOLATILE ORGANIC COMPOUNDS BY: SW 846 METHOD 8270 – Modified for Selected Ion Monitoring (SIM)

TABLE 6 (cont.)

## SVOA COMPOUNDS AND CHARACERISTIC IONS

2,4-Dinitrotoluene	165	63
4-Nitrophenol	109	139,65
2,3,4,6-Ttrachlorophenol	232	230
Diethylphthalate	149	177,176
Fluorene-d10 (surr)	176	174,178
Fluorene	166	165
4-Chlorophenyl-phenyl ether	204	206,141
4-Nitroaniline	138	108,65
4,6-Dinitro-2-methylphenol	198	121
N-nitrosodiphenylamine	169	168,167
4-Bromophenyl-phenyl ether	248	250,141
Hexachlorobenzene	284	142,249
Atrazine	200	173,215
Pentachlorophenol	266	264,268
Phenanthrene-d10 (IS)	188	189
Phenanthrene	178	179,176
Anthracene	178	179,176
Carbazole	167	166,139
Di-n-butylphthalate	149	150,104
Fluoranthene	202	200,203
Pyrene	202	200,201
Pyrene-d10 (surr)	212	210,106
Butylbenzylphthalate	149	91,206
Benzo(a)anthracene	228	229,226
Chrysene-d12 (IS)	240	236,120
3,3-Dichlorobenzidine	252	254,126
Chrysene	228	226,229
bis(2-Ethylhexyl)phthalate	149	167
Di-n-octylphthalate	149	150
Benzo(b)fluoranthene	252	253,125
Benzo(k)fluoranthene	252	253,125
Benzo(a)pyrene	252	253,250
Perylene-d12 (IS)	264	260
Indeno(1,2,3-cd)pyrene	276	277
Dibenzo(a,h)anthracene	278	279
Benzo(g,h,i)perylene	276	277

Primary ions must not be changed except in unusual instances where interference occurs with a co-eluting non-target analyte. In this case, a secondary ion may be used for quantitation with the following rules:

The quantitation ion must then be changed back to the one specified in the table above after quantitation of the samples(s).

Secondary ions are recommended only and may be changed depending upon instrument conditions (sensitivity, etc.). However, it is Katahdin policy that a minimum of 2 ions (primary and one secondary) be used for all GC/MS analyses.

<sup>(1)</sup> The corresponding standard(s) (initial calibration curve and continuing calibration standard) must be re-quantitated with the secondary ion.

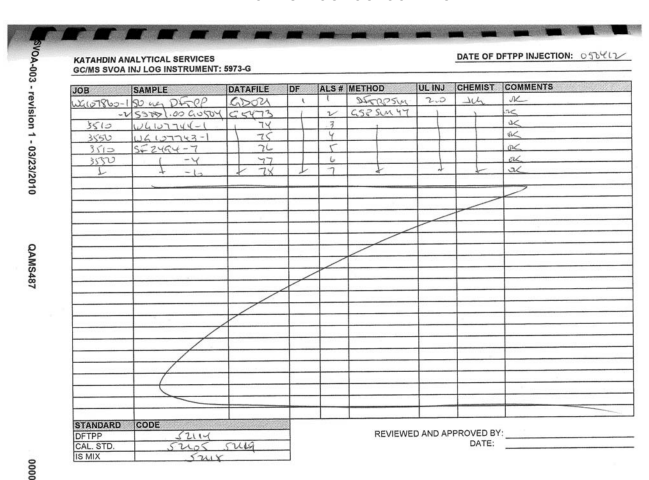
<sup>(2)</sup> Approval must be obtained from the Organic Department Manager or the laboratory Operations Manager.

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# TITLE: ANALYSIS OF SEMIVOLATILE ORGANIC COMPOUNDS BY: SW 846 METHOD 8270 – Modified for Selected Ion Monitoring (SIM)

FIGURE 1

EXAMPLE OF RUNLOG LOGBOOK PAGE



0000045

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#### TITLE: ANALYSIS OF SEMIVOLATILE ORGANIC COMPOUNDS BY: SW 846 METHOD 8270 - Modified for Selected Ion Monitoring (SIM)

### FIGURE 2

## EXAMPLE OF GC/MS STANDARDS RECEIPT LOGBOOK ENTRY

### KATAHDIN ANALYTICAL SERVICES

STOCK STANDARDS RECEIVED

GCMS LABORATORY REVIEWED BY/DATE: Anpoqub AccuStandard\* 125 Market St. Tel. 203 786 5 APP-9-176-D-20X
Pentachlorophenol
2.0 mg/mL in CH2Ct2
Lot: B3010100
Exp. Jan 10, 2013 Venl 3/16/01 ♠ AccuStandard\* FOR LABORATORY USE ONLY AMP2947 APP-9-090-50X 4,6-Dinitro-o-cresol 5.0 mg/mL in MeOH Lot: B1100296 Exp. Aug 16, 2012 ♠ AccuStandard® APP-9-145-50X
p-Nitrophenol
5.0 mg/mL in MeOH
Lot: B5050205
Exp. May 18, 2015 Ameggex QAMS294

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# TITLE: ANALYSIS OF SEMIVOLATILE ORGANIC COMPOUNDS BY: SW 846 METHOD 8270 – Modified for Selected Ion Monitoring (SIM)

FIGURE 3

EXAMPLE OF SVOA STANDARDS PREPARATION LOGBOOK ENTRY

	the addresses							Section and	Tooliyal	i Pinasona,
50863	8270 Stock	3-15-06	7-7-66	sin	Anpost4	8270 heyakus	300	222-07	4. aul	150 haghel
	(who ments)				ANPORST		350	3-15-67		0
			, was a same	C.M. P.S.	AMPO911	AP\$ 1x # 2	600	3-2-57		
		7 4 25			Amporio	+ 1	100	3-9-07		
					Ang ostu	1 1	200	7-7-06		
					Angolf9	organizatios pest	300	8-19-06		
					Anpoan	Beyon And		3-9-67		
					Anpoin			1.22.07	1,501	
					Anggok			3-9-67		
					Amp 536	3,3'- Dichlow bande	1	3-14-07		
					ANO932		150	3-9-04		
				1000000	50861	DEA	300	3-13-07		
		-			B13890	Mells	550			
50864	8270 level 1	3-15.06	7-7-06	الد	50863	800 Strile	70	7-7-06	1.05 ml	10 mglul
					B43590	rella	980			0
50865	8270 level 2	3-15-06	7-7-06	باند	50863	8270 Stock	150	7-7-06	vigonl	zoughel
					B43890	Melle	750			34
50866	8070 level 3	3-15-06	7-7-06	يالر	50X 3	suro Stale	600	7-706	1.8ml	50 mgho
					843690	mear	1200			
50867	8270 level 4	3-15-06	7-7-66	يار_	SUBLY	8270 Stak	700	7-7-06	1:05 ml	100 yell
					643890	Nelly	350			9

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# TITLE: ANALYSIS OF SEMIVOLATILE ORGANIC COMPOUNDS BY: SW 846 METHOD 8270 – Modified for Selected Ion Monitoring (SIM)

#### ADDENDUM 1

## LOW LEVEL 1,4-DIOXANE ANALYSIS

The following are differences from the standard 8270 C or D SIM analysis:

## GC Operating Conditions -

GC/MS Operating Conditions – Calibration and Samples			
Column temperature program	35°C for 4 min. to 300°C for 12°/min.		
Final column temperature hold	300°C		
	32 minutes (time may vary dependent upon column		
Run time	length)		
	2.0-3.0 minutes (time may vary dependent upon column		
Scan start time	length)		
Sample volume	2 uL		

Stock Standards – 1,4-Dioxane and 1,4-Dioxane each at 20 ug/mL

Calibration Standards – Use the above stock standards to prepare calibration standards at concentrations 0.25 ug/mL, 0.50 ug/mL, 1.0 ug/mL, 2.0 ug/mL, 4.0 ug/mL and 6.0 ug/mL. The 1.0 ug/mL is also the continuing calibration verification standard.

Sample analysis – Add 1 uL of internal standard (Section 5.3.2.4) aliquot of sample.

The ions for 1,4-Dioxane are 96 and 64.

# KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-515 Revision History Cover Page Page 1

TITLE: PREPARATION OF AQUEOUS SAMPLES FOR PESTICIDES/PCBs ANALYSIS				
Prepared By:	Mike Thomas	Date:	8/96	
Approved By:				
Group Supervisor:	michael F. Thomas	Date:	11/15/00	
Operations Manager:	\cBuston	Date:	10/25100	
QA Officer:	Detorah J. nadeau	Date:	10.23.00	
General Manager:	Dune f. hugan	Date:	11/16/00	
	,			
Revision History:				

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
01	Minor changes throughout. Clarifications to procedure Section.	9n	10:23:00	
02	Addition of SPE Propedure. Minor changes through out Addedwording to Sections 6 and 8	LAN	01810	013105
03	Added Separate Oc for Pest. and PCB. updated concentration procedure to re- flect current practices, Changes in wordin for clarification. Update Logbook page	LAO	04/06	04/06
04	Added waste generated and disposal info. Added missing definitions. Updated SPE extraction procedure. Updated Table land 2. Added Table 3.	LAD	०५।०७	०९१०७
05	opdated logbook example. Added logbook requirements	UAD	09/08	09/08

SOP Number: CA-515 Revision History Cover Page (cont.) Page 2

TITLE:	PREPARATION OF	AQUEOUS S	SAMPLES FOR	PESTICIDES/PCBs	<b>ANALYSIS</b>

Revision History:

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
06	Added information for determining initial volume. Added reference to CA-108. Added Clarification for LCS/D and MS/B Sets for PEST/PCB analysis. Minor changes to reflect current techniques.	ian)	10/09	10/09
70	Added additional solvent exchange procedure. Updated Logbook example.	LAY	08110	08/10
08	Added that a ms/msp should be extracted if enough sample volume and to extract an LCSD if no ms/D. Add wording to HSD4 pup, Minor changes to reflect current practices and remove duplication updated MDL-sect. 9. Added and updated references, Removed method 353 throughout. Changed PCB H.T to 30 Days w/explan	100	04/12	oulia
09	Sect. 7- Removed the procedure of menting the sample meniscus on the sample bothe with a grease pencil	LAO	05/13	05/13
	•			

Date Issued: 05/13

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TITLE:	PREPARATION OF AQUEOUS SAMPLES FOR PESTICIDES/PCBs ANALYSIS
	acknowledge receipt of this standard operating procedure by signing and dating both of the provided. Return the bottom half of this sheet to the QA Department.
l acknov	wledge receipt of copy of document SOP CA-515-09, titled PREPARATION OF US SAMPLES FOR PESTICIDES/PCBs ANALYSIS.
Recipien	nt: Date:
	DIN ANALYTICAL SERVICES, INC. ARD OPERATING PROCEDURE
	wledge receipt of copy of document SOP CA-515-09, titled PREPARATION OF US SAMPLES FOR PESTICIDES/PCBs ANALYSIS.
Recipien	nt:Date:

Date Issued: 05/13 Page 4 of 20

TITLE: PREPARATION OF AQUEOUS SAMPLES FOR PESTICIDES/PCBs ANALYSIS

### 1.0 SCOPE AND APPLICATION

The purpose of this SOP is to describe the procedures utilized by Katahdin Analytical Services, Inc. laboratory personnel for the preparation of aqueous samples prior to analysis for pesticides/PCBs by GC/ECD. It includes extraction of water samples by separatory funnel and continuous liquid-liquid extraction methods (EPA Methods 3510 and 3520.

#### 1.1 Definitions

METHOD BLANK (laboratory reagent blank): An artificial sample designed to determine if method analytes or other interferences are present in the laboratory environment, the reagents, or the apparatus. For aqueous samples, reagent water is used as a blank matrix; for soil samples, baked organic-free sand is used as a blank matrix. The blank is taken through the appropriate steps of the process.

LABORATORY CONTROL SAMPLE (LCS): A blank that has been spiked with the analyte(s) from an independent source, and is analyzed exactly like a sample. Its purpose is to determine whether the methodology is in control, and whether the laboratory is capable of making accurate and precise measurements. The matrix used should be phase matched with the samples and well characterized.

MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD): Predetermined quantities of stock solutions of certain analytes are added to a sample matrix prior to sample extraction and analysis. Samples are split into duplicates, spiked and analyzed. Percent recoveries are calculated for each of the analytes detected. The relative percent difference between the samples is calculated and used to assess analytical precision.

SURROGATES: Organic compounds which are similar to analytes of interest in chemical composition, extraction and chromatography, but which are not normally found in environmental samples. These compounds are spiked into all blanks, standards, samples and spiked samples prior to analysis. Percent recoveries are calculated for each surrogate.

### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of, analysts experienced in the extraction of aqueous samples for pesticides/PCBs analysis. Each analyst must demonstrate the ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, current revision, "Personnel Training".

It is the responsibility of all Katahdin personnel involved in the preparation of aqueous samples for pesticides/PCBs analysis to read and understand this SOP, adhere to the procedures outlined, and to properly document their data in the appropriate lab notebook. Any deviations from the test or irregularities with the

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#### TITLE: PREPARATION OF AQUEOUS SAMPLES FOR PESTICIDES/PCBs ANALYSIS

samples should also be recorded in the lab notebook and reported to the Department Manager or designated qualified data reviewer responsible for the data.

It is the responsibility of the Department Manager to oversee that members of their group follow this SOP, that their work is properly documented and to initiate periodic review of the associated logbooks.

## 1.3 Safety

Users of this procedure must be cognizant of inherent laboratory hazards, proper disposal procedures for contaminated materials and appropriate segregation of hazardous wastes. The toxicity or carcinogenicity of each reagent used in this method has not been precisely defined; however, each chemical should be treated as a potential health hazard. A reference file of material safety data sheets is available to all personnel involved in the chemical analysis. Everyone involved with the procedure must be familiar with the MSDSs for all the materials used in this procedure.

Each qualified analyst or technician must be familiar with Katahdin Analytical Environmental Health and Safety Manual including the Katahdin Hazardous Waste Management Plan and must follow appropriate procedures. These include the use of appropriate personal protective equipment (PPE) such as safety glasses, gloves and lab coats when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location of all safety equipment. Each analyst shall receive a safety orientation from their supervisor, or designee, appropriate for the job functions they will perform.

### 1.4 Waste Disposal

Wastes generated during the preparation of samples must be disposed of in accordance with the procedures described in the current revision of the Katahdin Hazardous Waste Management Plan and Safety Manual and SOP SD-903, "Sample Disposal," current revision. Expired standards are lab packed, placed in the Katahdin hazardous waste storage area, and disposed of in accordance with this SOP.

Any methylene chloride solvent waste generated during the rinsing of glassware etc. should be disposed of in the "D" waste stream satellite accumulation area nearest the point of generation. This includes the methylene chloride waste layer generated during CLLE extraction. Special care should be taken to pour this layer off into the appropriate waste stream, leaving the sample waste to be disposed of as follows. Since Pesticide/PCB samples are at a neutral pH, SEP funnel or CLLE sample waste may be dumped into either the "N-Hi" or "N-low" satellite accumulation area. Acetone and hexane are considered flammable waste, and should be disposed of in the "O" waste stream satellite accumulation area nearest the point of generation. Acid waste generated during the cleanup of PCB samples should be disposed of in the "O" satellite accumulation area nearest the point of generation. Please refer to

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TITLE: PREPARATION OF AQUEOUS SAMPLES FOR PESTICIDES/PCBs ANALYSIS

the current revision of SOP CA-107 for the location of satellite waste accumulation areas

### 2.0 SUMMARY OF METHOD

Pesticides/PCBs are extracted from aqueous samples using methylene chloride and separatory funnel or continuous liquid-liquid apparatus, following EPA Methods 3510 and 3520. The methylene chloride is exchanged with hexane for the final extract. Method detection limit studies must be performed annually for pesticides/PCBs using all extraction methods, if the extraction lab wishes to use either or all techniques. Method 3510 (separatory funnel) is generally preferred for pesticides/PCBs since organochlorine pesticides may dechlorinate if under elevated pH conditions for an extended period of time. (Section 3.2, Method 3510B, Rev. 2, 9/94)

#### 3.0 INTERFERENCES

Solvents, reagents, glassware, and other sample preparation apparatus may yield interferences to GC analysis due to the presence of contaminants. These contaminants can lead to discrete artifacts or elevated baselines in chromatograms. Routinely, all of these materials must be demonstrated to be free from interferences under the conditions of the analysis by running reagent blanks. Interferences caused by phthalate esters can pose a major problem in pesticide analysis. Common flexible plastics contain varying amounts of phthalates which are easily extracted during laboratory operations, so cross-contamination of glassware frequently occurs when plastics are handled. Interferences from phthalates can best be minimized by avoiding the use of such plastics in the laboratory. At no time may gloves which have not been tested for phthalates or gloves known to contain phthalates be used or stored in the organic extraction lab. Additionally, whenever possible plastic items in this lab must be replaced with metal or Teflon or other non-phthalate plastic substitute.

Special care should be taken to ensure clean glassware and apparatus are used, pre-rinsed with the appropriate solvent prior to use. Solvents should be analyzed prior to use to demonstrate that each lot is free of contaminants that may interfere with the analysis.

Interferences coextracted from the samples will vary considerably from source to source. If analysis of an extracted sample is prevented due to interferences, further cleanup of the sample extract may be needed to minimize interferences.

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#### TITLE: PREPARATION OF AQUEOUS SAMPLES FOR PESTICIDES/PCBs ANALYSIS

### 4.0 APPARATUS AND MATERIALS

Prior to use, all glassware must be rinsed three times with the solvent to be used for extraction.

- 4.1 Separatory Funnel 2000 mL capacity, Nalgene Teflon FEP separatory funnels with Nalgene Tefzel® screw-cap closures (or equivalent)
- 4.2 Concentrator tube 10 mL, graduated
- 4.3 Evaporative flask Kuderna-Danish, 500 mL capacity attached to concentrator with neck clips
- 4.4 Snyder column Kuderna-Danish, three ball macro
- 4.5 Graduated cylinders 100 mL, 1000 mL, or 2000 mL
- 4.6 Short Stem Funnels
- 4.7 250 mL amber collection bottles with Teflon-lined caps
- 4.8 12 mL and/or 16 mL glass vials with Teflon-lined caps
- 4.9 Continuous liquid-liquid extractors (CLLE) including body, 500 mL flat bottom boiling flask and Alhin condensers
- 4.10 Filter paper, 18.5 cm, Fisherbrand or Whatman 114V (or equivalent)
- 4.11 Nitrogen evaporation apparatus.
- 4.12 Boiling chips approximately 10/40 mesh, Teflon or selenized carborundum, 12 mesh (or equivalent). Cleaned by Soxhlet.
- 4.13 Water bath eight position concentric ring bath or equivalent, equipped with a calibrated thermometer.

### 5.0 REAGENTS

- 5.1 Laboratory reagent grade water water in which an interferent is not observed at or above the PQL for any parameter of interest (carbon filtered ASTM Type II water or equivalent)
- 5.2 Sodium Hydroxide (10N) Purchased from vendor, "Baker-analyzed", or equivalent

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#### TITLE: PREPARATION OF AQUEOUS SAMPLES FOR PESTICIDES/PCBs ANALYSIS

- 5.3 Sodium Sulfate (ACS) Granular, anhydrous. Bake at 400°C for 4 hours (may be done by vendor). Purify by rinsing three times with pesticide grade methylene chloride. Allow residual methylene chloride to evaporate before use. Stored in a Teflon capped glass bottle.
- 5.4 Sulfuric acid solution (1:1 H<sub>2</sub>SO<sub>4</sub>: H<sub>2</sub>O) Prepared in an icebath by slowly adding a volume of concentrated H<sub>2</sub>SO<sub>4</sub> to an equivalent volume of reagent water and swirl gently to mix. Caution should be taken when adding the acid to the water as the reaction is highly exothermic. Prepare as needed and store in a ground glass stoppered bottle.
- 5.5 Methylene Chloride (MeCL<sub>2</sub>) Pesticide grade or better. Lot must be verified by concentrating 300-400 mL to 1.0 mL and evaluating by GC/MS.
- 5.6 Acetone and Hexane Pesticide grade or better. Lot must be verified by concentrating approximately 20-30 mL to 1.0 mL and evaluating by GC/ECD.
- 5.7 Pesticide/PCB Surrogate spiking solution Prepare a solution of decachlorbiphenyl (DCB) and tetrachloro-meta-xylene (TCMX) at a concentration of 1.0 ug/mL ea in acetone. Store the solution at -10 to -20 °C in a Teflon sealed container. Solution must be verified by GC/ECD prior to use, and must be replaced every 6 months or sooner if degradation is evident.
- 5.8 Pesticide Matrix Spike/Lab Control Sample spiking solution Prepare a matrix spiking solution in pesticide grade methanol that contains all target analytes listed below:

ANALYTE	ug/mL
4,4'-DDT	0.5
4,4'-DDD	0.5
4,4'-DDE	0.5
Aldrin	0.5
Dieldrin	0.5
Endrin	0.5
Endrin Aldehyde	0.5
Endrin Ketone	0.5
Endosulfan I	0.5
Endosulfan II	0.5
Endosulfan Sulfate	0.5
alpha-BHC	0.5
beta-BHC	0.5
delta-BHC	0.5
gamma-BHC (Lindane)	0.5
Heptachlor	0.5
Heptachlor epoxide	0.5
Methoxychlor	0.5
alpha-Chlordane	0.5
gamma-Chlordane	0.5

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#### TITLE: PREPARATION OF AQUEOUS SAMPLES FOR PESTICIDES/PCBs ANALYSIS

5.9 PCB Matrix Spike/Lab Control Sample spiking solution - Prepare a matrix spiking solution in pesticide grade acetone that contains 5.0ug/ml ea of Aroclor® 1016/1260 mix (Restek catalog# 32039).

5.10 Store the spiking solutions at -10 to -20 °C in a Teflon sealed container. The solutions must be verified by GC/ECD prior to use, and must be replaced every 6 months or sooner if degradation is evident.

## 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

Samples are collected in 1 L amber bottles and held at 4 ( $\pm$ 2) °C until time of extraction.

The holding time for the extraction of aqueous pesticide samples for Methods 3510 and 3520 is 7 days from date of sample collection.

The holding time for the extraction of aqueous PCB samples by methods 3510 and 3520 is 30 Days

Note: SW-846 8082A does not specify a holding time to extraction; 30 days is presented here as a conservative limit. The method recommends a holding time of 40 days from extraction to analysis for extracts stored under refrigeration in the dark; but also refers to SW846 Chapter 4, which specifies that there is no holding time for PCBs. Additionally, SW-846 states that the holding times listed in the method under the conditions listed (apparently referring to storage of extracts) may be as long as a year.

Holding times may be dictated by a project specific Quality Assurance Project Plan (QAPP), in a program specific Quality Systems Manual (QSM) or by a regulating body. If a project requires a holding time that is not specified above it must noted in the analysis notes of the workorder.

#### 7.0 PROCEDURES

The following information must be recorded in the extraction logbook.

- Extraction method
- Surrogate and spike IDs
- Lot numbers of all solvents, acids and bases, sodium sulfate, filter paper
- Nitrogen evaporation water bath temperature
- Sample pH if applicable
- Extraction and Concentration dates
- Extraction and Concentration analyst
- Sample ID or QC sample ID
- Initial and final volumes or weight

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#### TITLE: PREPARATION OF AQUEOUS SAMPLES FOR PESTICIDES/PCBs ANALYSIS

- Surrogate and spike amounts
- Any sample cleanup preformed
- Final extract tray location
- Any comments regarding the sample extraction (ie. Emulsion)
- Prep batch start time and end time
- CLLE start time and end time

### SEPARATORY FUNNEL SAMPLE EXTRACTION

If an emulsion prevents acceptable recovery or client history indicates samples may demonstrate matrix interence, then samples should be extracted by continuous liquid-liquid extraction (CLLE).

- 7.1 Prerinse all glassware three times with methylene chloride prior to use.
- 7.2 Label a 2 L Teflon separatory funnel and a 250 mL amber collection bottle clearly. Label should include laboratory sample number, matrix, analyte, and extraction date. Be sure that the detachable stopcocks are secured to the separatory funnels before adding samples.
- 7.3 Measure the initial volume by comparing the meniscus of the sample with the reference bottle of the same bottle type. Please refer to SOP CA-108, "Basic Laboratory Technique", for the reference bottle verification procedure. Record the volume and any notable characteristics (e.g. color, presence of sediment, or odor) in the extraction logbook. Transfer the contents of the sample bottle to a 2 L separatory funnel.
- 7.4 Transfer 1 L of laboratory reagent grade water to a 2 L separatory funnel. This serves as a method blank for the extraction batch. A method blank must be prepared for every daily extraction batch of twenty or fewer samples.
- 7.5 Transfer 1 L of laboratory reagent grade water to a 2 L separatory funnel for each analysis to be performed (pesticide and/or PCB). This will serve as a Laboratory Control Sample (LCS). When Pesticides and PCBs are extracted together, a LCS and LCSD set must be extracted for each analysis. An LCS is required for every daily extraction batch of twenty or fewer samples and each analysis. If an MS/MSD pair is not extracted on a particular day, an LCS/LCSD pair may be required in order to meet client-specific or program-specific requirements. This information will be disseminated from the project manager or Department Manager.
- 7.6 A matrix spike/matrix spike duplicate (MS/MSD) is to be prepared as requested by a client or, at a minimum, one pair per 20 samples. An MS/MSD will be analyzed only if enough sample has been provided by the client. Additionally, in the event that the batch MS/MSD requirement cannot be fulfilled, a laboratory control spike duplicate must be analyzed. Transfer two additional 1 L aliquots of sample to 2 L separatory

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funnels for a matrix spike and matrix spike duplicate (MS/MSD) for each analysis. When Pesticides and PCBs are extracted together, a MS and MSD set must be extracted for each analysis. Note: Sufficient sample volume should be available without depleting all remaining sample aliquots.

- 7.7 Check the pH of the samples. If it is not between pH 5 and 9, adjust the pH with 10N sodium hydroxide or 1:1 sulfuric acid solution. Note the addition of NaOH or H<sub>2</sub>SO<sub>4</sub> in the extraction logbook.
- 7.8 Using a gas-tight syringe, add 1.0 mL of surrogate spiking solution to all samples the blank, LCS/LCSD(s) and MS/MSD(s), if performed.
- 7.9 Using a gas-tight syringe, add 1.0 mL of pesticide or PCB matrix spiking solution to the appropriate LCS, LCSD, MS and MSD if performed.
- 7.10 To each empty sample bottle add 60 mLs of methylene chloride, rinse the bottle and transfer the solvent into the appropriate separatory funnel. Add 60 mL of methylene chloride directly to the blank and LCS/LCSD(s).
- 7.11 Ensure that each screw cap is secured tightly to the separatory funnel to prevent leaks. Shake briefly and vent in hood to release pressure. Extract the sample by shaking the funnel on mechanical shaker for 3 minutes. Allow phases to separate for at least 10 minutes. Drain the methylene chloride layer into the 250 mL amber collection bottle.
- 7.12 If an emulsion forms, mechanical techniques must be employed to achieve maximum separation and solvent recovery. Such means include swirling and centrifugation and draining through a small separatory funnel. In certain instances, transferring the entire sample into a continuous liquid-liquid extractor may be the only alternative. If any such techniques are used, they must be noted in the extractions logbook.
- 7.13 Add a second 60 mL aliquot of methylene chloride to the separatory funnel and extract for the second time (see 7.11 7.13). Collect the methylene chloride layer in the same 250 mL amber collection bottle.
- 7.14 Repeat the extraction for a third time as described in 7.11 7.13.
- 7.15 Proceed to Section 7.30 for extract concentration procedures.

### CONTINUOUS LIQUID-LIQUID SAMPLE EXTRACTION (CLLE)

7.16 Set up the CLLE apparatus. All glassware should be rinsed three times with methylene chloride and the extract flasks properly labeled.

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7.17 Add 2-3 boiling stones to the round bottom flask and approximately 500 - 600 mL of methylene chloride to the CLLE body.

- 7.18 Add 1 L laboratory reagent grade water to a CLLE body. This is the method blank for this extraction batch. Be sure that no water leaks into the round bottom flask. A method blank is required for every extraction batch of twenty or fewer samples.
- 7.19 Measure the initial volume by comparing the meniscus of the sample with the reference bottle of the same bottle type. Please refer to SOP CA-108, "Basic Laboratory Technique", for the reference bottle verification procedure. Record the volume and any notable characteristics (e.g. color, presence of sediment, or odor) in the extraction logbook. Transfer the sample to a CLLE body, being sure that no water leaks into the round bottom flask.
- 7.20 Prepare an LCS for every daily extraction batch of twenty or fewer samples and each analysis (pesticide and/or PCB). Add 1 L of laboratory reagent grade water to a CLLE body. If an MS/MSD pair is not extracted on a particular day, an LCS/LCSD pair may be required in order to meet client-specific or program-specific requirements. This information will be disseminated from the project manager or Department Manager. When Pesticides and PCBs are extracted together, a LCS and LCSD set must be extracted for each analysis.
- 7.21 Check the pH of the samples. If it is not between pH 5 and 9, adjust the pH with 10N sodium hydroxide or 1:1 sulfuric acid solution. Note the addition of NaOH or H<sub>2</sub>SO<sub>4</sub> in the extraction logbook.
- 7.22 Transfer two 1 L portions of a sample to CLLE bodies for each analysis for preparation of a matrix spike/matrix spike duplicate if required. An MS/MSD is required if requested by the client or per 20 samples, whichever occurs first. When Pesticides and PCBs are extracted together, a MS and MSD set must be extracted for each analysis. Note: Sufficient sample volume should be available without depleting all remaining sample aliquots. An MS/MSD will be analyzed only if enough sample has been provided by the client. Additionally, in the event that the batch MS/MSD requirement cannot be fulfilled, a laboratory control spike duplicate must be analyzed.
- 7.23 For each sample, rinse the original sample container with approximately 30 mL of methylene chloride. Add this rinse to the CLLE body.
- 7.24 Add 1.0 mL of the Pesticide/PCB Surrogate Spike to each sample including the blank, LCS/LCSD and MS/MSD, if performed.
- 7.25 Add 1.0 mL of Pesticide or PCB Matrix Spike to the appropriate LCS/LCSD and MS/MSD pair, if performed, and stir.

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7.26 Attach cooling water Allihn condensers, after first rinsing each 45/50 joint with methylene chloride. Turn on the heating mantles and allow the samples to extract for at least 18 hours, total extract time may go up to 20 hours. Turn off the mantles and let samples cool.

7.27 Proceed to Section 7. 30 for sample extract concentration procedures.

#### CONCENTRATION OF WATER SAMPLE EXTRACTS

- 7.28 Rinse the K-D glassware (flask, concentration tube, funnel and Snyder column, including the ground glass joints on the flask and columns) three times with methylene chloride (or hexane for samples extracted with the Autoextractor) before assembling. Add two boiling chips to the K-D. Insert fluted 18.5 cm filter papers into short stem powder funnels and add ~ 2 inches of sodium sulfate crystals. Rinse the sodium sulfate in the assembled funnels with ~20-30 mLs of methylene chloride (hexane for samples extracted with the Autoextractor). Place the assembled K-D's under the funnels.
- 7.29 For methylene chloride extracts, add approximately 50 mL Hexane to funnel and let drain through. Since methylene chloride has a lower boiling point than Hexane, this will result in a final extract in hexane only.

**Note:** For Pesticide / PCB samples originating from South Carolina (see worknotes) do not add the hexane at this step. Solvent exchange will be during the nitrogen blow down procedure.

- 7.30 Transfer the methylene chloride or hexane extracts to the K-D concentrator setups through the sodium sulfate in the funnels. This is the drying step, which is required to remove residual water from the extracts. Any large water layers must be removed by other means, prior to pouring through the sodium sulfate. After pouring all of the extract volume through the sodium sulfate, rinse the extract bottle three times with  $\sim 2-3$  mLs of methylene chloride (or hexane for samples extracted with the Autoextractor). Add the rinsings through the sodium sulfate to complete a quantitative transfer. Rinse the sodium sulfate with  $\sim 15$  mLs of methylene chloride (or hexane for samples extracted with the Autoextractor) and allow to drain.
- 7.31 Transfer the labels from the collection bottles or round bottom flasks (from the CLLE extraction) to the K-Ds. Remove the funnel and attach a 3-ball macro Snyder column. Pre-wet the Snyder column with 1 mL of methylene chloride (or hexane for samples extracted with the Autoextractor).
- 7.32 Place the K-D in a hot water bath (85-90°C). Gently swirl K-D in the water until boiling begins. At the proper distillation rate, the Snyder balls should chatter but the chambers should not flood with condensed solvent. The K-D should be kept in a vertical orientation while on the bath. When the apparent volume in the concentrator tube reaches ≈ 5-6 mL, remove the K-D from the water bath. Allow the

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K-D to cool for 10 minutes. Rinse the Snyder column lower joint with  $\approx$  1 mL of hexane (methylene chloride for samples that have not gone through solvent exchange (ie. South Carolina samples). Remove the Snyder column. Wipe off any water from the neck above the lower joint of the flask. Separate the K-D flask from the concentrator tube, rinsing the ground glass joint with  $\approx$  1 mL hexane.

- 7.33 Reduce the extracts to ≈ 1 mL using Nitrogen blow-down apparatus. The bath temperature must be no higher than the boiling point of the solvent (45 °C for hexane). Turn the gas to 3 psi. Be careful not to splash the extract out of the tube. During concentration on the N-evap, the internal wall of the concentrator tube and the N-evap sparging pipet must be rinsed down at least once or twice with ≈1 mL of hexane (methylene chloride for samples not yet solvent exchanged). The solvent level in the concentrator tube must be positioned below the level of the water bath as much as possible to prevent water from condensing into the sample extract. As the extract volume is reduced, lower the N₂ sparging needle closer to the surface of the extract to expedite the concentration. Note any problems or extract losses, if they occur, in the extractions logbook. Transfer extract to a 12 or 16 mL vial. Using a reference vial for volume comparison, adjust the final extract volume to 10 mL by rinsing sides of tube with hexane and transferring rinsings to vial.
- 7.34 For samples that still need to be solvent exchanged, reduce the methylene chloride extract to ~ 1 mL. Add 10 mL of hexane to the concentrator tube and reduce to ~ 1 mL again on the N-evap. Adjust final extract to 10 mL by rinsing sides of tube with hexane and transferring rinsings to vial.
- 7.35 If at any point in the concentration procedure the concentrator tube goes dry reextract the sample immediately.
- 7.36 Transfer the label from the concentrator tube to the vial. Store extract vials at a temperature of  $4 \pm 2$  °C until ready for analysis. Indicate in the extractions logbook the box number and "tray location" of the individual extract vials.
- 7.37 All sample extracts for 8082 PCB analysis must undergo a sulfuric acid wash (cleanup) prior to analysis. All sample extracts for 8081 pesticide analysis do not undergo further cleanup unless requested by the client. Therefore, all sample extracts for combined 8081/8082 analyses must be split. Prior to splitting, mix contents of vial well. One portion must be acid cleaned for 8082 analysis. The associated method blank must be split and acid-cleaned in the same fashion. PCB LCSs and matrix spikes are acid cleaned also. Pesticide LCSs and matrix spikes are not subjected to further cleanup. Please refer to Katahdin SOP CA525 (current revision), Extract Cleanup Using Sulfuric Acid, for further instructions.

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### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

Each extractions analyst must demonstrate proficiency in performing the extractions that prepare samples for analysis. Demonstration consists of preparation (extraction), by the analyst, of at least four aliquots which are then analyzed according to the analytical method in question. These QC samples must meet all quality control acceptance limits. Demonstration must be documented by use of a form which summarizes the results of the analysis of these aliquots, calculated percent recoveries, and standard deviation. Demonstration of proficiency must be done one time per analyst initially and then annually thereafter. Refer to SOPs QA-805 and QA-807, current revision.

If, upon analysis of the extracted samples, it is discovered that quality control acceptance criteria have not been met, all associated samples must be evaluated against all the QC. In some cases data may be reported, perhaps with narration, while in other cases, other corrective action may be taken. The corrective actions may include re-extraction of the samples associated with the quality control sample that did not meet acceptance criteria, or may include making new reagents and standards if the standardization is suspect. These decisions are based on holding time considerations, client and project specific Data Quality Objectives and on review of chromatograms. The supervisor, Operations Manager, and/or Quality Assurance Officer may be consulted to evaluate data. Some samples may not be able to be reanalyzed within hold time. In these cases "qualified" data with narration may be advisable after consultation with the client.

Much of the work performed at the lab is analyzed in accordance with specific QC requirements spelled out in a project specific Quality Assurance Project Plan (QAPP) or in a program specific Quality Systems Manual (QSM). The reporting limits, acceptance criteria and/or corrective actions may be different than those specified in this SOP. In these cases the appropriate information will be communicated to the Department Manager and/or senior chemists before initiation of the analyses so that specific product codes can be produced for the project. In addition, the work order notes for each project will describe the specific QAPP or QSM to be followed.

A method blank must be extracted for each and every item listed below:

- Each day of extraction (24 hours midnight midnight)
- Each extraction method
- Every 20 samples extracted in a 24-hour period

A laboratory control sample (LCS) is required for each and every item listed below:

- Each extraction method
- Every extraction batch of twenty or fewer samples
- Each analysis (pesticide and/or PCB) to be performed

Refer to the current revision of the applicable Katahdin SOP for analysis of Pesticides and PCBs for quality control acceptance criteria.

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### 9.0 METHOD PERFORMANCE

The method detection limit (MDL) is defined as the minimum concentration of a substance that can be measured and reported with 99% confidence that the value is above zero. The MDLs shall be determined and verified one time per type of instrument unless otherwise required by the method.

A Limit of Detection (LOD) is an estimate of the minimum amount of a substance that an analytical process can reliably detect. An LOD is analyte and matrix specific and may be laboratory-dependent. LODs must be determined for all parameters for which the laboratory is accredited under the DoD Environmental Laboratory Accreditation Program. LOD's must be verified for every preparation and analytical method combination and on every applicable instrument on a quarterly basis.

The Limit of Quantitation (LOQ) is the minimum levels, concentrations, or quantities of a target variable (e.g., target analyte) that can be reported with a specified degree of confidence. The LOQ shall be set at the lowest point in the calibration curve for all analyses utilizing an initial calibration. LOQ's must be verified quarterly for every preparation and analytical method combination and on every applicable instrument on a quarterly basis for all parameters included in the DoD Scope of Accreditation. The LOQ must be verified at least once annually if the analysis is not included in the DoD Scope of Accreditation.

MDLs are filed with the Organic Department Manager and then with the QAO. LOD and LOQ verifications are filed with the QAO

Refer to the current revision of Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, for procedures on determining the MDL.

Refer to the current revision of the analytical SOP for other method performance parameters and requirements.

#### 10.0 APPLICABLE DOCUMENTS/REFERENCES

Test Methods for Evaluating Solid Waste - Physical/Chemical Methods, Methods 3510C and 3520C, USEPA SW-846, Third Edition, Final Update III, December 1996.

40 CFR 136, Appendix A, "Test Procedures for Analysis of Organic Pollutants," Method 608, June, 1998 edition.

The NELAC Institute, Laboratory Accreditation Standards, Volume 1, Management and Technical Requirements for Laboratories Performing Environmental Analysis, 10/06/2010.

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# TITLE: PREPARATION OF AQUEOUS SAMPLES FOR PESTICIDES/PCBs ANALYSIS

Department of Defense Quality Systems Manual for Environmental Laboratories (DOD QSM), Current Version.

# LIST OF TABLES AND FIGURES

Table 1	Summary of Method Modifications (Method 3510, Current Revision)
Table 2	Summary of Method Modifications (Method 3520, Current Revision)
Figure 1	Example of Runlog Page

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# TABLE 1 SUMMARY OF METHOD MODIFICATIONS (METHOD 3510, current revision)

TOPIC	KATAHDIN SOP CA-515-09	METHOD 3510, current revision
Apparatus/Materials  Reagents	<ol> <li>1. 12 or 16 mL vials used for final extract</li> <li>2. 250 mL amber bottle or flask used</li> <li>3. 1.0 mL syringe</li> <li>4. short stem funnels</li> </ol>	2 mL vials used for final extract     2. 250 mL Erlenmeyer flask     5.0 mL syringe     drying column
Sample preservation/ handling	entire contents of 1 L sample bottle transferred to separatory funnel	one liter graduated cylinders used to transfer initial sample volume to separatory funnel
Procedures	<ol> <li>extract collection in amber bottle or Erlenmeyer flask</li> <li>extract dried using Na<sub>2</sub>SO<sub>4</sub> in short stem funnels</li> <li>no apparatus height specification for concentration on water bath</li> <li>sample removed from water bath when volume reaches ~10 mL</li> <li>hexane added directly to K-D body at start of concentration process (this modification is not allowed for samples originating from South Carolina).</li> </ol>	<ol> <li>extract collection in Erlenmeyer flask</li> <li>extract dried using Na<sub>2</sub>SO<sub>4</sub> in drying columns</li> <li>partially immerse concentrator tube in water and lower apparatus to complete concentration in 10-15min</li> <li>sample removed from water bath when volume reaches 1-2 mL</li> <li>solvent exchange via large K-D with addition of 50 mL hexane after concentrating methylene chloride extract to 1 mL</li> </ol>
QC - Spikes		
QC - LCS		
QC - Accuracy/Precision		
QC - MDL		

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# TITLE: PREPARATION OF AQUEOUS SAMPLES FOR PESTICIDES/PCBs ANALYSIS

# TABLE 2 SUMMARY OF METHOD MODIFICATIONS (METHOD 3520, current revision)

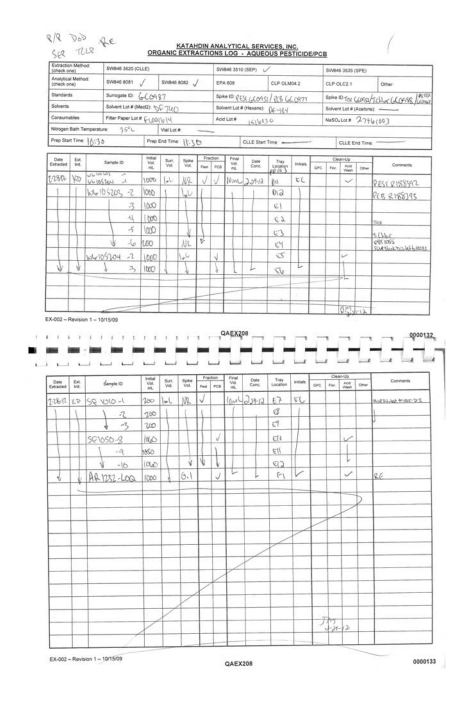
TOPIC	KATAHDIN SOP CA-515-09	METHOD 3520, current revision
Apparatus/Materials	<ol> <li>short-stem funnels</li> <li>12 or 16 mL vials used for final extract</li> </ol>	drying columns     2 mL vials used for final extract
Reagents		
Sample preservation/ handling	entire contents of 1 L sample bottle transferred to CLLE	one liter graduated cylinders used to tranfer initial sample volume to CLLE
Procedures	<ol> <li>CLLE for 18 ± 2 hours</li> <li>extract dried using Na<sub>2</sub>SO<sub>4</sub> in short stem funnels</li> <li>no apparatus height specification for concentration on water bath</li> <li>sample removed from water bath when volume reaches ~10 mL</li> <li>hexane added directly to K-D body at start of concentration process (this modification is not allowed for samples originating from South Carolina).</li> </ol>	<ol> <li>CLLE for 18-24 hours</li> <li>extract dried using Na<sub>2</sub>SO<sub>4</sub> in drying columns</li> <li>partially immerse concentrator tube in water and lower apparatus to complete concentration in 10-15min</li> <li>sample removed from water bath when volume reaches 1-2 mL</li> <li>solvent exchange via macro K-D with addition of 50 mL hexane after concentrating methylene chloride extract to 1 mL</li> </ol>
QC - Spikes	,	
QC - LCS		
QC - Accuracy/Precision		
QC - MDL		

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FIGURE 1
EXAMPLE OF LOGBOOK PAGE



## KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-524 Revision History Cover Page Page 1

	TION OF SEDIMENT/SOIL SAMPLES BY SETHOD 3540 FOR PESTICIDE/PCB ANALYS		EXTRACTION
Prepared By:	Mike Thomas	Date:	7/98
Approved By:			
Group Supervisor:	Michael F. Thomas	Date:	11/15/00
Operations Manager:	\ Benta	Date:	11/15/00
QA Officer:	Queborah J. nadean	Date:	11.15.00
General Manager:	Deman P. hufah	Date:	11/16/00
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Revision History:			

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
01	Minor changes throughout. Clarifications to procedure Section.	<i>9</i> n	11:15:00	11/13/00
02	Added definitions to section 1.1. wording changed or added to clarify sections 5, 6, 8, +9. New figure		11.08.04	11, 08,04
03	Sect. 7.1.2 - adding the step to rinse forceps also. 7.10 adding condenser temperature and output voltage of verriable transformer	LAD	04/06	04/06
04	Added generated weste information. Updated spike list. Added LCSD. Reworded Sect. 7.10 and 7.11 for clarification. Updated Table! Replaced Figure 1	LAD	09/07	०९/०७
05	Changed "N. Lo" waste to "K" waste. Updated Logbook example. Sect. 7-added wording instructing the recording of consumable lot #5 in Logbook.	UAID	07/08	07/08

SOP Number: CA-524 Revision History Cover Page (cont.) Page 2

TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
06	Added balance criteria. Changed weight criteria to 7-0.059. Minor changes to section? to reflect current techniques. Clarified samples being GPC'd are not solvent exchanged into hexare. Updated logbook page. Added CA-108 reference for subsempling information	LAD	08109	90180
07	Removed taugeting sample weights. Removed decanting samples prior to extraction.	LAD	08/10	08/10
08	minor changes to section 7 to reflect Current practices. Updated Section 7.6 for trequency of MS/D's. Added information for MDL, LODand LOQ to Section. 9. Updated	LAD	04/12	04/12
	for MDL, LODand LOQ to Section. 9. Updated references. Added the Soy to reasents.			
				71111

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TITLE:	PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS
Please a spaces p	acknowledge receipt of this standard operating procedure by signing and dating both of the provided. Return the bottom half of this sheet to the QA Department.
SEDIME	vledge receipt of copy of document SOP CA-524-08, titled PREPARATION OF ENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR IDE/PCB ANALYSIS.
Recipien	nt:Date:
	DIN ANALYTICAL SERVICES, INC. ARD OPERATING PROCEDURE
SEDIME	viedge receipt of copy of document SOP CA-524-08, titled PREPARATION OF ENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR IDE/PCB ANALYSIS.
Recipien	nt:Date:

SOP Number: CA-524-08

Date Issued: 04/12

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### TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

#### 1.0 SCOPE AND APPLICATION

The purpose of this SOP is to describe the procedure for extracting pesticides/PCBs from solids such as soils, sludges, and wastes using Method 3540. The Soxhlet extraction process ensures intimate contact of the sample matrix with the extraction solvent.

This method is applicable to the isolation and concentration of water-insoluble and slightly water-soluble organics in preparation for a variety of chromatographic procedures including methods 8081 for pesticides and 8082 for PCB's.

#### 1.1 Definitions

METHOD BLANK (laboratory reagent blank): An artificial sample designed to determine if method analytes or other interferences are present in the laboratory environment, the reagents, or the apparatus. For aqueous samples, reagent water is used as a blank matrix; for soil samples, baked organic-free sand is used as a blank matrix. The blank is taken through the appropriate steps of the process.

LABORATORY CONTROL SAMPLE (LCS): A blank that has been spiked with the analyte(s) from an independent source, and is analyzed exactly like a sample. Its purpose is to determine whether the methodology is in control, and whether the laboratory is capable of making accurate and precise measurements. The matrix used should be phase matched with the samples and well characterized.

MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD): Predetermined quantities of stock solutions of certain analytes are added to a sample matrix prior to sample extraction and analysis. Samples are split into duplicates, spiked and analyzed. Percent recoveries are calculated for each of the analytes detected. The relative percent difference between the samples is calculated and used to assess analytical precision.

SURROGATES: Organic compounds which are similar to analytes of interest in chemical composition, extraction and chromatography, but which are not normally found in environmental samples. These compounds are spiked into all blanks, standards, samples and spiked samples prior to analysis. Percent recoveries are calculated for each surrogate.

#### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of analysts experienced in the extraction of samples for pesticide/PCB analysis. Each analyst must demonstrate and document their ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, current revision, "Personnel Training and Documentation of Capability".

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

It is the responsibility of all Katahdin technical personnel involved in the extraction of samples for pesticide/PCB analysis to read and understand this SOP, adhere to the procedures outlined, and to properly document their data in the appropriate lab notebook. Any deviations from the test or irregularities with the samples should also be recorded in the lab notebook and reported to the Department Manager or designated qualified data reviewer responsible for this data.

It is the responsibility of the Department Manager to oversee that members of their group follow this SOP, to ensure that their work is properly documented and to indicate periodic review of the associated logbooks.

#### 1.3 Safety

Users of this procedure must be cognizant of inherent laboratory hazards, proper disposal procedures for contaminated materials and appropriate segregation of hazardous wastes. The toxicity or carcinogenicity of each reagent used in this method has not been precisely defined; however, each chemical should be treated as a potential health hazard. A reference file of material safety data sheets is available to all personnel involved in the chemical analysis. Everyone involved with the procedure must be familiar with the MSDSs for all the materials used in this procedure.

Each qualified analyst or technician must be familiar with Katahdin Analytical Environmental Health and Safety Manual including the Katahdin Hazardous Waste Plan and must follow appropriate procedures. These include the use of appropriate personal protective equipment (PPE) such as safety glasses, gloves and lab coats when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location of all safety equipment. Each analyst shall receive a safety orientation from their Department Manager, or designee, appropriate for the job functions they will perform.

#### 1.4 Waste Disposal

Wastes generated during the preparation of samples must be disposed of in accordance with the procedures described in the current revision of the Katahdin Hazardous Waste Plan and Safety Manual and SOP SD-903, "Sample Disposal," current revision. Expired standards are lab packed, placed in the Katahdin hazardous waste storage area, and disposed of in accordance with this SOP.

Any methylene chloride solvent waste generated during the rinsing of glassware etc. should be disposed of in the "D" waste stream satellite accumulation area nearest the point of generation. Acetone and hexane are considered flammable waste, and should be disposed of in the "O" waste stream satellite accumulation area nearest the point of generation. Post-extraction soil samples, used glass wool, and sodium sulfate waste should be disposed of in the soil with organics "I" waste stream

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

satellite accumulation area nearest the point of generation. Acid waste generated during the cleanup of PCB samples should be disposed of in the "K" satellite accumulation area nearest the point of generation. Please refer to the current revision of SOP CA-107 for the location of satellite waste accumulation areas.

#### 2.0 SUMMARY OF METHOD

- 2.1 The solid sample is mixed with anhydrous sodium sulfate, placed in a Soxhlet extractor and extracted with methylene chloride.
- 2.2 The extract is then dried, concentrated, and exchanged into hexane for GC analysis. Sulfuric acid cleanup is performed on extracts for 8082 PCB analysis.

#### 3.0 INTERFERENCES

Solvents, reagents, glassware, and other sample preparation apparatus may yield interferences to GC analysis due to the presence of contaminants. These contaminants can lead to discrete artifacts or elevated baselines in chromatograms. Routinely, all of these materials must be demonstrated to be free from interferences under the conditions of the analysis by running reagent blanks. Interferences caused by phthalate esters can pose a major problem in pesticide analysis. Common flexible plastics contain varying amounts of phthalates that are easily extracted during laboratory operations, so cross-contamination of glassware frequently occurs when plastics are handled. Interferences from phthalates can best be minimized by avoiding the use of such plastics in the laboratory. At no time may gloves that have not been tested for phthalates or gloves known to contain phthalates be used or stored in the organic extraction lab. Additionally, whenever possible plastic items in this lab must be replaced with metal or teflon or other non-phthalate plastic substitute.

Special care should be taken to ensure clean glassware and apparatus are used, prerinsed with the appropriate solvent prior to use. Solvents should be analyzed prior to use to demonstrate that each lot is free of contaminants that may interfere with the analysis.

Interferences coextracted from the samples will vary considerably from source to source. If analysis of an extracted sample is prevented due to inteferences, further cleanup of the sample extract may be needed to minimize interferences.

#### 4.0 APPARATUS AND MATERIALS

- 4.1 Soxhlet extractor 45/50 top joint and 24/40 lower joint.
  - 4.1.1 500 mL flat-bottom boiling flask

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

- 4.1.2 Allihn cooling water condenser
- 4.2 Powder Funnels 100 mm top diameter, 35 mm stem
- 4.3 Kuderna-Danish (K-D) apparatus
  - 4.3.1 Concentrator tube 10-mL
  - 4.3.2 Evaporation flask 500-mL
  - 4.3.3 Snyder column Three-ball macro
- 4.4 Nitrogen evaporation (N-EVAP) apparatus.
- 4.5 Boiling stones, 12 mesh silicon carbide (carborundum) pre-purified by Soxhlet extraction in methylene chloride
- 4.6 Water bath Heated, with concentric ring cover, capable of temperature control (± 5°C). The bath should be used in a hood.
- 4.7 Vials Glass, 4, 12, or 16 mL with Teflon-lined screw caps
- 4.8 Glass wool (fiberglass) baked at 400°C for a minimum of 4 hours or overnight.
- 4.9 Heating mantles Rheostat controlled.
- 4.10 Disposable glass Pasteur pipets, 5 ¾", and bulbs.
- 4.11 Drying oven capable of maintaining 105°C for glassware drying.
- 4.12 Muffle oven capable of maintaining 400 °C for baking glass wool and organic-free sand.
- 4.13 Beakers, 250 or 400 mL
- 4.14 Top-loading balance capable of weighing to 0.01 g.
- 4.15 Spatulas, stainless-steel
- 4.16 Long forceps, stainless-steel
- 4.17 Metal clips for securing Soxhlets to boiling flasks
- 4.18 Filter paper, 18.5 cm, Fisherbrand or Whatman 114V (or equivalent)

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

4.19 Gel Permeation Chromatograph (GPC) - J2 Scientific AccuPrep MPS<sup>™</sup> with internal UV detection

#### 5.0 REAGENTS

- 5.1 Sodium sulfate (granular, anhydrous and powdered, anhydrous) (ACS reagent grade), Na<sub>2</sub>SO<sub>4</sub>. Certified by the manufacturer/vendor as purified by heating at 400°C for 4 hours prior to receipt by the laboratory.
- 5.2 Sulfuric acid solution (1:1 H2SO4 : H2O) Prepared in an icebath by slowly adding a volume of concentrated H2SO4 to an equivalent volume of reagent water and swirl gently to mix. Caution should be taken when adding the acid to the water as the reaction is highly exothermic.
- 5.3 Methylene chloride (pesticide grade or equivalent) purchased by lot, evaluated prior to use by concentration of 300 mL to 1 mL followed by GC/MS analysis.
- 5.4 Acetone and hexane (pesticide grade or equivalent) purchased by lot, evaluated prior to use by concentration of 300 mL to 1 mL followed by GC/MS and GC analysis.
- 5.5 Organic-free sand, purified by baking at 400 °C at a minimum of 4 hours or overnight. Method blanks serve as checks on the baked sand.
- 5.6 Surrogate spiking solution Prepare a solution of decachlorobiphenyl (DCB) and tetrachloro-meta-xylene (TCMX) at a concentration of 1 ug/mL in acetone.
- 5.7 Matrix Spike/Lab Control Sample spiking solution
  - 5.6.1 Pesticide spike solution prepare in pesticide grade methanol containing the analytes listed below at concentrations of 0.5 ug/mL.

4,4'-DDD
4,4'-DDE
4,4'-DDT
Aldrin
alpha-BHC
beta-BHC
delta-BHC
Dieldrin
Endosulfan I
Endosulfan Sulfate

Endrin
Endrin Aldehyde
Endrin Ketone
gamma-BHC (Lindane)
Heptachlor
Heptachlor Epoxide
Methoxychlor
alpha-Chlordane
gamma-chlordane
Endrin
Endrin Aldehyde

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

- 5.6.2 PCB spike solution prepare Aroclor 1660 (Aroclor 1016 and 1260) in pesticide grade acetone at a concentration of 5.0 ug/mL each.
- 5.7 Store the solutions mentioned in sections 5.5 and 5.6 at -10 to -20 °C (±2 °C) in a Teflon sealed container. Solution must be verified by GC/ECD prior to use and must be replaced every 6 months or sooner if degradation is evident.

#### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

Sediment/soil samples must be collected in a soil jar and must be maintained at 4°C (±2°C).

Holding time for extraction of sediment/soil samples for Method 3540 is 14 days from date of sample collection, although the analyst should be aware that actual holding times employed may be project/program specific.

Store all extracts at 4°C (±2°C) in the dark in labeled Teflon-sealed containers. See SOP SD-902, "Sample Receipt and Internal Control," current revision, for storage areas and temperature maintenance procedures.

#### 7.0 PROCEDURES

The following information must be recorded in the extraction logbook.

- Extraction method
- Surrogate and spike IDs
- Lot numbers of all solvents, acids and bases, sodium sulfate, filter paper
- Nitrogen evaporation water bath temperature
- pH if applicable
- Extraction and Concentration dates
- Extraction and Concentration analyst
- Soxhlett extraction start and end times, also the prep start and end times
- Sample ID or QC sample ID
- Initial and final volumes or weight
- Surrogate and spike amounts
- Final extract tray location
- Any comments regarding the sample extraction (ie. Emulsion)

Samples need to be "swiped" out when removing and "swiped" in when replacing samples in storage locations to maintain the internal chain of custody. Refer to Katahdin SOP, Sample Receipt and Internal Control, current revision, for the proper procedure for removal and return samples.

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

#### 7.1 Preparing the Soxhlet Extraction Apparatus

- 7.1.1 Rinse the Soxhlet extractors and 500 mL flat-bottom boiling flasks three times with methylene chloride. Be sure that the solvent rinses through the large vapor tube and smaller siphon tubes of the Soxhlet. Inspect these for tiny cracks. Also rinse the 24/40 lower joint.
- 7.1.2 Add ~ 250 mLs of methylene chloride to the 500 mL boiling flask. Add several boiling stones. Rinse the stainless steel forceps with Methylene chloride. Working in a hood, place a plug of the glass wool at the bottom of the Soxhlet so that the siphon tube hole is covered. Insert the 24/40 joint of the Soxhlet extractor into the 500 mL boiling flask and secure with a metal clip. Cover the top of the Soxhlet extractor with a piece of aluminum foil until ready to begin loading the sample.

#### 7.2 Sample Handling

- 7.2.1 Sediment/soil samples Do not decant any water layer on a sediment sample. Mix the sample thoroughly with the stainless steel spatula. If the sample container is full to the extent that stirring the sample is impractical, try to remove the "best representative" aliquot from the jar based on color, particle size, moisture, etc. Discard any foreign objects such as sticks, leaves, and rocks.
- 7.2.2 Gummy, fibrous, or oily materials not amenable to mixing should be cut, shredded, or otherwise reduced in size to allow for maximum exposure of the sample surfaces to the extraction solvent. Materials such as glass, rubber, metal, etc. may not require mixing with powdered sodium sulfate to disperse the sample. Plastic materials must be tested for degradation (melting) in methylene chloride prior to Soxhlet extraction.
- 7.2.3 Please refer to Katahdin Analytical Services SOP CA-108, current revision, "Basic Laboratory Techniques" for more information of subsampling.
- 7.3 Weigh out an approximate, greater than 30 g portion of sample into a labeled 400 mL beaker. Record sample weight to nearest 0.01 g in appropriate extraction logbook. Add between 30 to 60 g of powdered sodium sulfate, as required, to produce a "free-flowing" mixture. The amount of sodium sulfate added will depend upon the moisture content of the sample (e.g., low moisture content will require less sodium sulfate). Mix well with a spatula. Keep the spatula in the sample beaker and cover the beaker with aluminum foil. Record sodium sulfate lot in logbook.
- 7.4 A method blank must be prepared with each extraction batch, not to exceed 20 client samples. To prepare method blank, weigh out one greater than 30 g portion of purified

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

sand in a labeled 400 mL beaker. Record sample weights to nearest 0.01 g in appropriate extraction logbook. Add 60 g sodium sulfate and mix well. (Although a "free-flowing" mixture can be achieved with less than 60 g sodium sulfate, the method blank must contain 60 g in order to evaluate the sodium sulfate as a potential source of contamination.)

- 7.5 A laboratory control sample (LCS) must be prepared with each extraction batch, not to exceed 20 client samples. To prepare LCS, weigh out one greater than 30 g portion of purified sand in a labeled 400 mL beaker. Record sample weights to nearest 0.01 g in appropriate extraction logbook. Add 30 g sodium sulfate and mix well. With extraction batches prepared for combined 8081/8082 Pesticide and PCB analysis, separate Pesticide and PCB LCS's must be prepared (refer to section 5.6). If an MS/MSD pair is not extracted on a particular day, an LCS/LCSD pair may be required in order to meet client-specific or program-specific requirements. This information will be disseminated from the project manager or Department Manager.
- 7.6 A matrix spike/matrix spike duplicate (MS/MSD) set should be prepared for every 20 samples. An MS/MSD will be analyzed only if enough sample has been provided by the client. Additionally, in the event that the batch MS/MSD requirement cannot be fulfilled, a laboratory control spike duplicate must be analyzed. To prepare MS/MSD, weigh out two approximate, greater than 30g portions of the sample designated for MS/MSD into each of two labeled 400 mL beakers. Record sample weights to nearest 0.01g in appropriate extraction logbook. Add 30 g sodium sulfate to each to produce a free-flowing mixture, and mix well. With extraction batches prepared for combined 8081/8082 Pesticide and PCB analysis, separate Pesticide and PCB MS/MSD pairs must be prepared (refer to section 5.6).
- 7.7 Once all of the QC and field samples have been weighed and mixed with sodium sulfate, begin adding each to the assembled and appropriately labeled Soxhlet extractors using the stainless steel spatulas. Carefully scrape all of the mixtures from the beaker walls so that no more than 1% remains behind in the beaker. Be careful that none of solid material falls into the extract flask through the large vapor tube.
- 7.8 To all samples, method blank, LCS/LCSD, and MS/MSD add 1.0 mL (if FV=10mL, adjust amount for different final volumes) of the pesticide/PCBs surrogate spiking solution using a 1.0 mL gas tight syringe. Record surrogate spike volume and identification code in the extraction logbook. Thoroughly rinse syringe with solvent between each use.
- 7.9 To LCS/LCSD and MS/MSD add 1.0 mL (if FV=10mL, adjust amount for different final volumes) of either the pesticide or PCBs matrix spike/LCS spiking solutions using a 1.0 mL gas tight syringe. Record matrix spike/LCS spiking solution volume

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

and identification codes in the extraction logbook. Thoroughly rinse syringe with solvent between each use.

- 7.10 Rinse the joints of the Allihn cooling condensers with Methylene Chloride, collecting the waste in a methylene chloride solvent waste container. Place each of the Soxhlet extractors in a heating mantle and lower the Allihn cooling water condensers into the 45/50 joints of the extractors. The condensers should be set to a temperature of 15°C. Save the pieces of aluminum foil for covering the Soxhlets when the extraction is complete. Switch on the individual heating mantles and be sure that the Rheostat of the variable transformer is set to 55% of the output voltage. Once the methylene chloride begins to boil and the Soxhlet begins to cycle (solvent will immerse the sample and collect in the Soxhlet until the level reaches that of the small siphon tube and then begin to spill over into the extract flask), recheck the apparatus' for leaks. Allow the samples to extract for 18-24 hours. Be sure the chiller/recirculator temperature is set low enough to provide enough cooling capacity for the number of extractions in the batch.
- 7.11 When the extraction is complete, allow the extracts to cool before dismantling. Remove the Allihn condenser and replace the aluminum foil on top of the extractor. Move the extractors to a hood and detach the extractor from the extract flask. Tilt each extractor slightly to cause any remaining solvent in the sample chamber to drain through the siphon tube into the extract flask. This will help to cool the extract flask and make the apparatus easier to dismantle. Try to drain as much solvent as possible from the extractor into the flask. This is done by rinsing a glass tube in methylene chloride and pressing on the sample slightly so that as solvent as possible is drained into the extract flask. Cover the flask with aluminum foil and store in the interim extract storage refrigerator unless the extracts are to be concentrated the same day.
- 7.12 Immediately remove the extracted soil/sodium sulfate mixtures from the extractors using a square edge spatula, and dispose of in an appropriate solid waste container. It is important to do this soon after the extractors are dismantled, as the sample mixture will tend to "freeze" into a solid mass in the Soxhlet as the solvent dries.

#### CONCENTRATION OF THE EXTRACTS

7.13 Rinse the K-D glassware (flask, concentration tube, and Snyder column, including the ground glass joints on the flask and columns) three times with methylene chloride before assembling. Add a few boiling chips to the K-D. Insert 18.5 cm filter papers into short stem powder funnels and add ~ 2 inches of sodium sulfate crystals. Place the assembled K-D's under the funnels. Record the lot numbers of the solvent, sodium sulfate and filter papers in the extraction logbook.

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

- 7.14 If samples are to be GPC'd, refer to the current revision of Katahdin SOP CA-513, Extract Cleanup Using Gel Permeation Chromatography, for appropriate concentrating procedures. Samples that undergo GPC are not solvent exchanged into hexane. All pesticide soil samples should be cleaned up to reduce matrix interferences.
- 7.15 If samples are not to be GPC'd follow Steps 7.16 through 7.23 to concentrate extracts to final volume of 10 mLs (or a client specified final volume)
- 7.16 For a solvent exchange, (for samples not being GPC'd), add approximately 50 mL hexane to funnel and let drain through. Since methylene chloride has a lower boiling point than Hexane, this will result in a final extract in hexane only. Record the lot number of the solvent in the extraction logbook.
- 7.17 Transfer the extracts to the K-D concentrator setups through the sodium sulfate in the funnels. This is the drying step, which is required to remove residual water from the extracts. Any large water layers must be removed by other means, prior to pouring through the sodium sulfate. After pouring all of the extract volume through the sodium sulfate, rinse the extract flask three times with  $\sim 2-3$  mLs of methylene chloride. Add the rinsings through the sodium sulfate to complete a quantitative transfer. Rinse the sodium sulfate with  $\sim 15$  mLs of methylene chloride and allow to drain.
- 7.18 Transfer the labels from the extract flasks to the K-Ds. Remove the funnel and attach a 3-ball macro Snyder column. Pre-wet the Snyder column with 1 mL of methylene chloride.
- 7.19 Place the K-D in a hot water bath (75-85°C). Gently swirl K-D in the water until boiling begins. At the proper distillation rate, the Snyder balls should chatter but the chambers should not flood with condensed solvent. The K-D should be kept in a vertical orientation while on the bath. When the apparent volume in the concentrator tube reaches ≈ 6 mL, remove the K-D from the water bath. Allow the K-D to cool for 10 minutes. Rinse the Snyder column lower joint with ≈ 1 mL of methylene chloride, hexane, if exchange is taking place. Remove the Snyder column. Wipe off any water from the neck above the lower joint of the flask. Separate the K-D flask from the concentrator tube, rinsing the ground glass joint with ≈ 1 mL methylene chloride, hexane, if exchange is taking place.
- 7.20 Reduce the extract in the concentrator tube to approximately 1-2 mL using the nitrogen blow-down apparatus to ensure all methylene chloride has been evaporated. The bath temperature must be no higher than the boiling point of the solvent (39°C for methylene chloride). Turn the gas to 3 psi. Be careful not to splash the extract out of the tube. <u>During concentration on the N-evap</u>, the internal wall of the concentrator tube and the N-evap sparging pipet must be rinsed down at least

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

once or twice with  $\approx 1$  mL of hexane. The solvent level in the concentrator tube must be positioned below the level of the water bath as much as possible to prevent water from condensing into the sample extract. As the extract volume is reduced, lower the  $N_2$  sparging pipet closer to the surface of the extract to expedite the concentration. Record the temperature of the water in the nitrogen evaporation water bath in the extraction logbook, also note any problems or extract losses, if they occur.

- 7.21 Complete quantitative transfer of the extract to a vial by using hexane. Adjust the volume of the hexane extract to 10 mL (or a client specified final volume) in either a 12 or 16 mL vial using the appropriate "reference vial" for volume comparison.
- 7.22 Label the vial with lab sample number, extraction date, matrix and analysis. Store extract vials at a temperature of  $4 \pm 2$  °C until ready for analysis. Indicate in the extractions logbook the box number and "tray location" of the individual extract vials.
- 7.23 All sample extracts for 8082 PCB analysis must undergo a sulfuric acid wash (cleanup) prior to analysis, unless it has been GPC'd. All sample extracts for 8081 pesticide analysis should undergo further cleanup using the GPC unless time is a factor. All sample extracts for combined 8081/8082 analyses must be split unless GPC'd. One portion must be acid cleaned for 8082 analysis. The associated method blank must be split and acid-cleaned in the same fashion. PCB LCSs and matrix spikes are acid cleaned also. Pesticide LCSs and matrix spikes are not subjected to further cleanup. Record the lot number of the acid in the extraction logbook. Please refer to Katahdin SOP CA525 (current revision), Extract Cleanup Using Sulfuric Acid, for further instructions.

#### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

A method blank must be extracted for <u>each</u> and <u>every</u> item listed below:

- Each sample matrix (soil, water)
- Each day of extraction (24 hours midnight midnight)
- Each extraction method or level
- Every 20 samples extracted in a 24-hour period

A laboratory control sample (LCS) is required for <u>each</u> and <u>every</u> item listed below:

- Each sample matrix
- Each extraction method or level
- Every extraction batch of twenty or fewer samples
- Each analysis (pesticide and/or PCB) to be performed

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

Refer to the current revision of the applicable Katahdin SOP for analysis of Pesticides and PCBs for quality control acceptance criteria.

Each extractions analyst must demonstrate proficiency in performing the extractions that prepare samples for analysis. Demonstration consists of preparation (extraction), by the analyst, of at least four aliquots which are then analyzed according to the analytical method in question. These QC samples must meet all quality control acceptance limits. Demonstration must be documented by use of a form which summarizes the results of the analysis of these aliquots, calculated percent recoveries, and standard deviation. Demonstration of proficiency must be done one time per analyst initially and then annually thereafter. Refer to SOPs QA-805 and QA-807, current revision.

If, upon analysis of the extracted samples, it is discovered that quality control acceptance criteria have not been met, all associated samples must be evaluated against all the QC. In some cases data may be reported, perhaps with narration, while in other cases, other corrective action may be taken. The corrective actions may include re-extraction of the samples associated with the quality control sample that did not meet acceptance criteria, or may include making new reagents and standards if the standardization is suspect. These decisions are based on holding time considerations, client and project specific Data Quality Objectives and on review of chromatograms. The supervisor, Operations Manager, and/or Quality Assurance Officer may be consulted to evaluate data. Some samples may not be able to be reanalyzed within hold time. In these cases "qualified" data with narration may be advisable after consultation with the client.

Much of the work performed at the lab is analyzed in accordance with specific QC requirements spelled out in a project specific Quality Assurance Project Plan (QAPP) or in a program specific Quality Systems Manual (QSM). The reporting limits, acceptance criteria and/or corrective actions may be different than those specified in this SOP. In these cases the appropriate information will be communicated to the Department Manager and/or senior chemists before initiation of the analyses so that specific product codes can be produced for the project. In addition, the work order notes for each project will describe the specific QAPP or QSM to be followed.

#### 9.0 METHOD PERFORMANCE

The method detection limit (MDL) is defined as the minimum concentration of a substance that can be measured and reported with 99% confidence that the value is above zero. The MDLs shall be determined and verified one time per type of instrument unless otherwise required by the method.

A Limit of Detection (LOD) is an estimate of the minimum amount of a substance that an analytical process can reliably detect. An LOD is analyte and matrix specific and may be laboratory-dependent. LODs must be determined for all parameters for which the laboratory

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

is accredited under the DoD Environmental Laboratory Accreditation Program. LOD's must be verified for every preparation and analytical method combination and on every applicable instrument on a quarterly basis.

The Limit of Detection (LOQ) is the minimum levels, concentrations, or quantities of a target variable (e.g., target analyte) that can be reported with a specified degree of confidence. The LOQ shall be set at the lowest point in the calibration curve for all analyses utilizing an initial calibration. LOQ's must be verified quarterly for every preparation and analytical method combination and on every applicable instrument on a quarterly basis for all parameters included in the DoD Scope of Accreditation. The LOQ must be verified at least once annually if the analysis is not included in the DoD Scope of Accreditation.

MDLs are filed with the Organic Department Manager and then with the QAO. LOD and LOQ verifications are filed with the QAO

Refer to the current revision of Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, for procedures on determining the MDL.

Refer to the applicable analytical SOP for other method performance parameters and requirements.

#### 10.0 APPLICABLE DOCUMENTS/REFERENCES

Test Methods for Evaluating Solid Waste-Physical/Chemical Methods, Method 3540C, SW-846, Third Edition, Updates I, II, IIA, IIB, and III Revised December 1996, US EPA.

Department of Defense Quality Systems Manual for Environmental Laboratories (DOD QSM), Current Version.

The NELAC Institute, Laboratory Accreditation Standards, Volume 1, Management and Technical Requirements for Laboratories Performing Environmental Analysis, 10/06/2010...

#### LIST OF TABLES AND FIGURES

Table 1 Summary of Method Modifications

Figure 1 Example of Logbook Page

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## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

## TABLE 1 SUMMARY OF METHOD MODIFICATIONS

TOPIC	KATAHDIN SOP CA-524-08	METHOD 3540, current revision
Apparatus/Materials	short stem funnels	1. drying columns
Reagents		
Sample preservation/ handling		
Procedures	<ol> <li>Use 30 grams of sample and 30 grams of sodium sulfate.</li> <li>Use 250 mL of methylene chloride</li> <li>no apparatus height specification for concentration on water bath</li> <li>water bath at 75-85 deg C</li> <li>sample removed from water bath when volume reaches ~6 mL</li> <li>Solvent exchange to hexane is performed using K-D apparatus with addition of approximately 50 mL hexane at the start of concentration process</li> </ol>	<ol> <li>Use 10 grams of sample and 10 grams of sodium sulfate.</li> <li>Use 300 mL of methylene chloride</li> <li>partially immerse concentrator tube in water and lower apparatus to complete concentration in 10-15min</li> <li>water bath at 80-90 deg C</li> <li>sample removed from water bath when volume reaches 1-2 mL</li> <li>Solvent exchange to hexane is performed using K-D apparatus with addition of approximately 50 mL hexane after concentrating methylene chloride extract to 1 mL</li> </ol>
QC - Spikes		
QC - LCS		
QC - Accuracy/Precision		
QC - MDL		

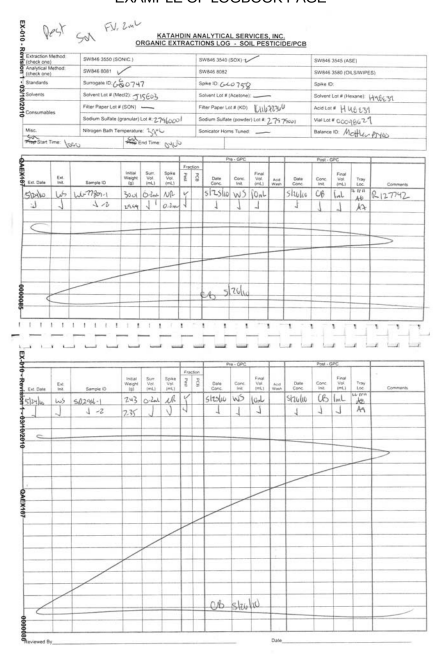
SOP Number: CA-524-08

Date Issued: 04/12 Page 18 of 18

## TITLE: PREPARATION OF SEDIMENT/SOIL SAMPLES BY SOXHLET EXTRACTION USING METHOD 3540 FOR PESTICIDE/PCB ANALYSIS

FIGURE 1

EXAMPLE OF LOGBOOK PAGE



# ADDENDUM SOP NO CHANGE FORM

## KATAHDIN ANALYTICAL SERVICES, INC. SOP "REVIEW WITH NO CHANGES" FORM

Name of Person Reviewing SOP: Jessica Spear	-in-Wildes
Review Date: 3-15-13	
SOP Number: CA-524	
· · · · · · · · · · · · · · · · · · ·	Isoil Samples by Soxhle
SOP Title: Preparation of Sedment extraction using method 3540	for Rest IPCBS
THE ABOVE REFERENCED SOP HAS BEEN REVIEWED ANALYST OR SUPERVISOR. NO CHANGES ARE REQUI	BY A QUALIFIED AND TRAINED
Department Supervisor Signature:	Date:
biter J	3-19-13
QAO Signature:	Date:
Lesein Dimond	032113

## KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-329 Revision History Cover Page Page 1

TITLE: ANALYSIS OF PCBs AS TOTAL AROCLORS BY GAS CHROMATOGRAPHY/ ELECTRON CAPTURE DETECTOR (GC/ECD): SW-846 METHOD 8082

Prepared By:	Peter Lemay	Date:_	4/98
Approved By:			
Group Supervisor:	Daten Len	Date:_	1/15/01
Operations Manager	: John C. Benton	Date:_	V15107
QA Officer:	O Deborah J. nadeau	Date:_	1.22.01
General Manager:	Deman P. Lukare	Date:_	1/16/01
Revision History:			1 . /

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
01	Format changes, added pollution prevention, minor changes to Sections 7,8 and Table 1	D	1-22-01	1/00/01
8082	TIONS 7, 8 WILL INDICE			
02	Revised Sections 7.3.1, 7.4.5 and 7.6.1 to be compliant with South Cardina	Dn	5.23.01	5.23.01
8082	requirements.			
03	Changed to practice of reporting higher value. Other minor changes	Dn	5.21.02	5.21.02
8082	to sections 7.5.2, 7.7.3 + to			
04	Revised SOP to indicate Turbochrom is			
8082	being used as instrument wontrol + data collection software. Included Target-re- lated definitions. Changes to sections 7.7.3, 7.7.4 and 7.8.	MRC	08.20.04	08.20.04
05	Changed 7.5.2 to reflect alternating CV			
8085	Changed Table 2 Sect. 7.3.1 New Checklist	LAD	020305	020305
	added wording to sect. 8			ANN

SOP Number: CA-329 Revision History Cover Page Page 2

TITLE: ANALYSIS OF PCBs AS TOTAL AROCLORS BY GAS CHROMATOGRAPHY/ ELECTRON CAPTURE DETECTOR (GC/ECD): SW-846 METHOD 8082

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
9087	Changed PCB 1260 to Aroclor 1260. Removed references to 3541. UPdated table 2. Added instructions to shake extract before vialing	LAD	04/06	04/06
רס	Added weste streams to sect. 1.0. Added ICV to definitions, sect. 5, sect. 7 and Table 1. Added wording regarding 2nd column confirmation criteria and flagging whesto sect. 7.7.4. Added CCV criteria to sect. 7.5.3 and Table 1. Added wording regarding MI to sect. 7.7.3	LAD	08/07	08/07
08	Added tissue, wipe and oil matrices. Added extraction method 3535. Added DDT anolog interference, Std. information and analysis prequency criteria. Added HTs are a recommendation. Added note that 2 detections to be used for aual column. Updated method references lemoved calibration and surrogate method mod. from Tab. 3. Added more into a linear calib. Added extraction	fors LA-V)	02/09	03/09
09	references, Added Chemstedion to definitions. Clerified that Surrogates are added to only the aroclor 1660 standards, not ALL standards.	LAN	05109	05109
10	Revised Sections 7, 8, and 10 to applicate compliance with the DOD QSM version 4.1	LAD	08/09	08/09
11	Added Table 2 with DOD QSM Ver. 4.1 QC criteria. Minor changes to Table 1.	(An	04/10	04/10
12	Removed Sect. 4.5- Analytical balance, Removed Sect. 5.24.  DUT analog standard lemoved Sect. 7.5- DUT analog  Standard consequirement. Table 1- Added aver. cel.  Criteria and consected ecv LCS acceptance criteria.  Added and removed referencesto Sect. 10. Updated  Figure 2-deta review checklish. Added PCBs 1262 2, 1268	LAO	07/11	07/4
13	Added Extraction method 3546. Removed Quickforms references. Added reporting from Kims. updated Figures 1 and 2	UAD	09/13	02/13

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#### 1.0 SCOPE AND APPLICATION

This SOP describes all aspects of the analysis of extracts of aqueous, solid, tissue, wipe and oil samples for PCBs by EPA Method 8082A as performed by Katahdin Analytical Services, Inc. including sample analysis, data review, standard preparation and instrument calibration.

It is applicable to the following compounds: Aroclor-1016, Aroclor-1221, Aroclor-1232, Aroclor-1242, Aroclor-1248, Aroclor-1254, Aroclor-1260, Aroclor-1262 and Aroclor-1268. Extracts are analyzed by Gas Chromatography-Electron Capture Detector (GC-ECD).

#### 1.1 Definitions

ANALYTICAL BATCH: 20 or fewer samples which are analyzed together with the same method sequence and the same lots of reagents and with the manipulations common to each sample within the same time period or in continuous sequential time periods.

METHOD BLANK (laboratory reagent blank): An artificial sample designed to determine if method analytes or other interferences are present in the laboratory environment, the reagents, or the apparatus. For aqueous samples, laboratory reagent grade water is used as a blank matrix; for soil samples, muffled sand is used as a blank matrix. The blank is taken through the appropriate steps of the process.

CALIBRATION CHECK: Verification of the ratio of instrument response to analyte amount, a calibration check is done by analyzing for analyte standards in an appropriate solvent. Calibration check solutions are made from a stock solution that is different from the stock used to prepare standards.

CALIBRATION STANDARD (WORKING STANDARD): A solution prepared from the stock standard solution that is used to calibrate the instrument response with respect to analyte concentration.

INDEPENDENT CALIBRATION VERIFICATION STANDARD (ICV): A solution prepared from a stock standard solution independent of the calibration mix that is used to verify the calibration.

LABORATORY CONTROL SAMPLE (LCS): A blank that has been spiked with the analyte(s) from an independent source and is analyzed exactly like a sample. Its purpose is to determine whether the methodology is in control, and whether the laboratory is capable of making accurate and precise measurements. The matrix used should be phase matched with the samples and well characterized.

MATRIX SPIKE/MATRIX SPIKE DUPLICATE (MS/MSD): Predetermined quantities of stock solutions of certain analytes are added to a sample matrix prior to sample extraction and analysis. Samples are split into duplicates, spiked and analyzed. Percent

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recoveries are calculated for each of the analytes detected. The relative percent difference between the samples is calculated and used to assess analytical precision.

STANDARD CURVE (CALIBRATION CURVE): A curve that plots concentration of known analyte standard versus the instrument response to the analyte.

STOCK STANDARD SOLUTION: A concentrated solution containing a single certified standard that is a method analyte, or a concentrated solution of a single analyte prepared in the laboratory with an assay reference compound. Stock standard solutions are used to prepare calibration standards.

SURROGATES: Organic compounds which are similar to analytes of interest in chemical composition, extraction and chromatography, but which are not normally found in environmental samples. These compounds are spiked into all blanks, Aroclor 1660 standard, samples and spiked samples prior to analysis. Percent recoveries are calculated for each surrogate.

KATAHDIN INFORMATION MANAGEMENT SYSTEM (KIMS): A complete multi-user system with the capabilities of integrating laboratory instrumentation, generating laboratory worksheets, providing complete Lab Order status and generating reports. KIMS utilizes these features through a database.

PE NELSON TURBOCHROM OR HP CHEMSTATION: data acquisition systems that are used to collect chromatographic data. The systems can also be used to archive raw data files.

TARGET: A software system that combines full processing, reporting and comprehensive review capabilities, regardless of chromatographic vendor and data type.

TARGET DB: An oracle database used to store and organize all Target data files.

#### 1.2 Responsibilities

- 1.2.1 This method is restricted to use by, or under the supervision of analysts experienced in the analysis of PCBs by method 8082. Each analyst must demonstrate and document their ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, "Personnel Training & Documentation of Capability," current revision.
- 1.2.2 It is the responsibility of all Katahdin technical personnel involved in analysis by method 8082 to read and understand this SOP, adhere to the procedures outlined, and to properly document their data in the appropriate lab notebook. Any deviations from the test or irregularities with the samples should also be

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recorded in the lab notebook and reported to the Department Manager or designated qualified data reviewer responsible for this data.

1.2.3 It is the responsibility of the Department Manager to oversee that members of their group follow this SOP, to ensure that their work is properly documented and to initiate periodic review of the associated logbooks.

#### 1.3 Health and Safety

- 1.3.1 Users of this procedure must be cognizant of inherent laboratory hazards, proper disposal procedures for contaminated materials and appropriate segregation of hazardous wastes. The toxicity or carcinogenicity of each reagent used in this method has not been precisely defined; however, each chemical should be treated as a potential health hazard. A reference file of material safety data sheets is available to all personnel involved in the chemical analysis. Everyone involved with the procedure must be familiar with the MSDSs (material safety data sheets) for all the materials used in this procedure.
- 1.3.2 Each qualified analyst or technician must be familiar with Katahdin Analytical Environmental Health and Safety Manual including the Katahdin Hazardous Waste Plan and must follow appropriate procedures. These include the use of appropriate personal protective equipment (PPE) such as safety glasses, gloves and lab coats when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location of all safety equipment. Each analyst shall receive a safety orientation from their Department Manager, or designee, appropriate for the job functions they will perform.

#### 1.4 Pollution Prevention/Waste Disposal

- 1.4.1 Whenever possible, laboratory personnel should use pollution prevention techniques to address their waste generation. Refer to the current revision of the Katahdin Hazardous Management Program for further details on pollution prevention techniques.
- 1.4.2 Wastes generated during the preparation of samples must be disposed of in accordance with the Katahdin Analytical Environmental Health and Safety Manual and SOP SD-903, "Sample Disposal," current revision. Expired standards are lab packed, placed in the Katahdin hazardous waste storage area, and disposed of in accordance with this SOP.
- 1.4.3 Wastes generated during standards preparation are disposed of in the Mixed Flammable Waste (O). After the extracts have been analyzed, the autosampler vials and any expired standard vials or ampules are disposed of in the PCB Vial Waste (H).

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#### 2.0 SUMMARY OF METHOD

- 2.1 Method 8082 provides gas chromatographic conditions for the detection of PPB concentrations of certain PCBs. Prior to the use of this method, appropriate sample extraction techniques must be used. Both neat and diluted organic liquids (Method 3580, waste dilution) may be analyzed by direct injection. A 2 to 5 ul aliquot of sample is injected into a gas chromatograph (GC) using the direct injection technique, and compounds in the GC effluent are detected by an electron capture detector (ECD).
- 2.2 The sensitivity of Method 8082 usually depends on the concentration of interferences rather than on instrumental limitations. If interferences prevent detection of the analytes, Method 8082 may also be performed on samples that have undergone the following cleanups: Method 3660 Sulfur Cleanup and Method 3665 Sulfuric Acid Cleanup.

#### 3.0 INTERFERENCES

Interferences by phthalate esters can pose a problem in PCB determinations when using the electron capture detector. Common flexible plastics contain various amounts of phthalates. Care has to be taken to avoid using any plastic materials during the extraction process. Exhaustive cleanup of reagents and glassware may be required to eliminate background phthalate contamination.

Compounds from the sample matrix to which the detector will respond, such as single-component chlorinated pesticides including the DDT series.

#### 4.0 APPARATUS AND MATERIALS

#### 4.1 Gas chromatograph

- 4.1.1 GC Hewlett Packard 5890 series I or II connected to the Turbochrom or HP Chemstation data system, or equivalent.
- 4.1.2 Columns Instruments are configured with a pre-column originating from the injection port, which is connected to a deactivated glass Y splitter that connects two different columns to two detectors. The most commonly used columns are: RTX-35 30M x 0.53 mm ID, RTX-5 30M x 0.53 MM ID, or RTX-1701 30M x 0.53 mm ID. Equivalent columns can be used.
- 4.1.3 Detectors: Electron capture detectors (ECD). Note: Two detectors must be employed when using dual columns.
- 4.2 Volumetric flasks, class A: sizes as appropriate with the ground-glass stoppers.

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- 4.3 Syringes: various sizes for preparing standards and injecting samples on the instrument.
- 4.4 Vials: various sizes and types including crimp tops.
- 4.5 Refrigerator for storage of extracts and standards.

#### 5.0 REAGENTS

- 5.1 Solvents
  - 5.1.1 Hexane: pesticide quality or equivalent for diluting samples and standards.
- 5.2 Standards
  - 5.2.1 Stock standard solutions: Solutions purchased from suppliers like Restek or other acceptable retailers. Expiration dates are one year from date of opening vial or sooner if manufacturers date is less. Upon receipt, all standards are logged into the appropriate logbook with the date of receipt, expiration date, source, lot number, solvent and concentration of compounds. Standard solutions are stored at 4°C in polytetrafluoroethylene (PTFE)-sealed containers in the dark.
  - 5.2.2 Calibration standards: Prepared through the dilution of the stock standards with hexane. Expiration date is 6 months or sooner. Information is documented in standards prep logbook. The concentrations of the working PCB calibration standards are 0.05 ug/ml, 0.10 ug/ml, 0.25 ug/ml, 1.0 ug/ml, 2.5 ug/ml, and 10.0 ug/ml. The Aroclor 1660 standard also contain the surrogates Tetrachloro-m-xylene (TCX) and Decachlorobiphenyl (DCB) at the respective concentrations: 0.001 ug/ml, 0.002 ug/ml, 0.005 ug/ml, 0.020 ug/ml, 0.050 ug/ml, and 0.20 ug/ml.
  - 5.2.3 Independent Calibration Verification standard (ICV): Prepared through the dilution of the stock standards with hexane. Expiration date is 6 months or sooner. Information is documented in standards prep logbook. The concentration of the ICV PCB standard is 1.0 ug/ml.

#### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

Extracts must be stored under refrigeration and analyzed within 40 days of extraction.

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**Note:** The holding time above is a recommendation. PCBs are very stable in a variety of matrices, and holding times under the conditions listed above may be as long as a year.

#### 7.0 PROCEDURES

#### 7.1 Extraction

Refer to the appropriate SOPs for the correct extraction procedure. In general, water samples are extracted using methods 3510, 3520 or 3535 while solid samples use methods 3540, 3545, 3546 or 3550. Tissue samples are extracted using method 3545 or 3540. Wipes and oils are generally extracted using method 3580.

#### 7.2 Instrument conditions

Refer to the instrument logbook for the current column and conditions.

#### Typical conditions are:

Makeup flow: 60 ml/min Helium, Ar/Methane or Nitrogen

Column flow: 6 ml/min Injector Temp: 200 Detector Temp: 300

Oven Ramp: 160(0) - 5/min - 260(10)

Run time: 30 min Injection size: 2 ul

#### 7.3 Calibration

7.3.1 The GC system is calibrated using the external standard calibration procedure. Six-point calibration standards of Aroclor 1660 (Aroclor 1016 and Aroclor 1260), Aroclor 1242, Aroclor 1248 and Aroclor 1254 are prepared. Six-point calibration standards of Aroclor 1221, Aroclor 1232, Aroclor 1262 and Aroclor 1268 are also prepared. If Aroclor 1221, Aroclor 1232, Aroclor 1262 and Aroclor 1268 are suspected, then a six-point curve of the respective Aroclor will be analyzed prior to the analysis of samples. At a minimum, a single point calibration standard is analyzed for these Aroclors. If using a single point and the Aroclor is required for a project and is detected in a sample, then the GC would be calibrated for the Aroclor and the samples would be reanalyzed.

Each calibration standard is injected using the technique that is used to introduce the actual samples into the GC. Three to five characteristic peaks from each Aroclor are used to calibrate a curve. The Target system will calculate a peak height for all three to five peaks in each Aroclor. A separate

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calibration curve for each of the three to five peaks can be prepared in Target using the peak height against the concentration of the standard. A non-linear calibration applying a second order polynomial (quadratic fit) equation is used to prepare the curve. In order to be used for quantitative purposes, the Coefficient of Determination (r²) must be greater than or equal to 0.990. The quadratic equation is:

 $y = ax^2 + bx + c$ 

where: y = Instrument response

b = Slope of the line

x = Concentration of the calibration standard

c = the intercept

- 7.3.2 A non-linear calibration model may not be allowable for certain states, federal programs, or clients. South Carolina does not allow non-linear calibration work originating in their state. In these cases, a linear calibration model must be used. Each calibration standard is injected using the technique that is used to introduce the actual samples into the GC. Three to five characteristic peaks from each Aroclor are used to calibrate a curve. The Target system will calculate a peak height for all three to five peaks in each Aroclor. A separate calibration curve for each of the three to five peaks can be prepared in Target using the peak height against the concentration of the standard.
  - 7.3.2.1 Linear calibration using the average calibration factor

The calibration factor (CF) is calculated using the following formula:

Where:  $A_s = Peak$  area (or height) of the analyte or surrogate.

 $C_s$  = Concentration of the analyte or surrogate, in  $\mu g/L$ .

To evaluate the linearity of the initial calibration, calculate the mean CF, the standard deviation (SD), and the RSD.

If the RSD of the calibration factor is less than or equal to 20% over the calibration range, then linearity through the origin may be assumed, and the average calibration or response factor may be used to determine sample concentrations.

7.3.2.2 Linear calibration using a least squares regression

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where: y = Instrument response

b = Slope of the line

x = Concentration of the calibration standard

c = the intercept

The analyst should not force the line through the origin, but have the intercept calculated from the five data points. In addition, do not include the origin (0,0) as a sixth calibration point. The regression calculation will generate a correlation coefficient (r) that is a measure of the "goodness of fit" of the regression line to the data. A value of 1.00 indicates a perfect fit. In order to be used for quantitative purposes, r must be greater than or equal to 0.995. The ICAL must be successful before any samples or other QC check samples can be analyzed.

- 7.3.3 The AR1660 calibration curve must be checked initially by analyzing a standard containing the same analytes as the curve but prepared from another source. If the response of the analytes from the independent source varies by more than  $\pm$  20%, a new independent source standard must be analyzed or a new calibration curve must be prepared and/or analyzed.
- 7.3.4 The working calibration curve must be verified prior to sample analysis and every 10 samples thereafter by injecting the mid-point calibration standard. If the response for any analyte varies from the expected response by more than  $\pm$  15%, a new calibration curve must be prepared for that analyte. The average result for 5 peak heights of the Aroclors is used for quantitation.

For clients or projects requiring DoD QSM 4.1, the response for any analyte must not vary from the expected response by more than  $\pm$  20%, or a new calibration curve must be prepared for that analyte. If the CCV fails the above criteria, reanalyze all samples since the last successful calibration verification. If reanalysis cannot be performed, data must be qualified and explained in the narrative. Additionally, apply a Q-flag to all results for the specific analyte(s) in all samples since the last acceptable calibration verification.

#### 7.4 Retention time windows

- 7.4.1 Three injections are made of all the PCBs throughout the course of a 72 hour period.
- 7.4.2 A major peak from the envelope is chosen and a standard deviation is calculated using the three retention times for that peak.

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7.4.3 Plus or minus three times the standard deviation of the retention times for each standard is used to define the retention time window; however, the experience of the analyst should weight heavily in the interpretation of chromatograms. The analyst should use the retention time window, but should primarily rely on pattern recognition.

- 7.4.4 Retention time windows are calculated for each standard on each GC column at method setup and after major maintenance, including whenever a new GC column is installed. The data is kept on file in the laboratory.
- 7.4.5 If the calculated retention time window results in a value of 0.03 minutes or less, the laboratory will apply nominal windows. This is done in order to avoid any false negative hits because of the window being to narrow. The windows are:  $\pm$  0.07 for all target analytes. By utilizing these windows, a false positive hit may be initially indicated, but an experienced analyst could determine a false positive from scrutinizing the chromatograms. Please note that the use of nominal retention time windows may not be allowable for certain states, federal programs, or clients. South Carolina does not allow the use of nominal limits for compliance work originating in their state. In these cases, a window of  $\pm$  0.03 minutes must be used if the established retention time window is less than 0.03 minutes.

#### 7.5 Gas chromatographic analysis

- 7.5.1 Shake samples and let them sit for one minute before vialing for analysis.
- 7.5.2 All instrument injections are performed using the direct injection technique with an autosampler set for 2-5 ul injection volumes.
- 7.5.3 Samples are analyzed in a set referred to as an analysis sequence. The sequence begins with instrument calibration as listed in section 7.3 followed by sample extracts interspersed with mid-concentration calibration standards. Before any samples are analyzed the instrument must be calibrated by analyzing a six-point calibration or a 1.0ppm concentration standard (CVcalibration verification standard) for Aroclor 1660, Aroclor 1242, Aroclor 1248 and Aroclor 1254. If a CV is run, the calculated concentration must not exceed a difference of  $\pm$  15%. For clients or projects requiring DoD QSM 4.1, the response for any analyte must not vary from the expected response by more than + 20%, or a new calibration curve must be prepared for that analyte. If the CCV fails the above criteria, reanalyze all samples since the last successful calibration verification. If reanalysis cannot be performed, data must be qualified and explained in the narrative. Additionally, apply a Q-flag to all results for the specific analyte(s) in all samples since the last acceptable calibration verification. Each sample analysis must be bracketed with an acceptable initial

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calibration or an opening CV and an ending CV for each 12-hour shift. The closing CV for Aroclor 1660 is a 0.25ppm concentration standard. All other Aroclors at the closing of the run remain at 1.0ppm concentration. If a second window of samples is run immediately after the closing CVs, the concentration of Aroclor 1660 at the completion of this window would be 1.0ppm. The calibration standard must also be injected at intervals of not less than once every ten samples and at the end of the analysis sequence. If the CV fails, the instrument is checked for any obvious problems and maintenance is performed if deemed necessary. Another CV is analyzed or the instrument is recalibrated and then samples are injected. All samples that were injected after the standard exceeding the criterion must be reinjected to avoid errors in quantitation, if the initial analysis indicated the presence of the specific target analyte that exceeded the criterion.

- 7.5.3.1 However, if the standard analyzed <u>after</u> a group of samples exhibits a response for an analyte that is <u>above</u> the acceptance limit, i.e. >15%, and the analyte was not detected in the specific samples analyzed during the analytical shift, then the extracts for those samples do not need to be reanalyzed, as the CV standard has demonstrated that the analyte would have been detected were it present. In contrast, if an analyte above the QC limits <u>was</u> detected in a sample extract, then reinjection is necessary to ensure accurate quantitation. If an analyte was not detected in the sample and the standard response is more than 15% below the initial calibration response, then re-injection is necessary to ensure that the detector response has not deteriorated to the point that the analyte would not have been detected even though it was present.
- 7.5.4 The center of the retention time window for each analyte and surrogate is established by using the absolute retention time for each analyte and surrogate from the daily opening calibration verification or initial calibration.
- 7.5.5 The identification of PCBs is based on agreement between the retention times of peaks in the sample chromatogram with the retention time windows established through the analysis of standards of the target analytes. An analyte is tentatively identified when a peak from a sample falls within the daily retention time window. Each tentative identification must be confirmed using a second GC column of dissimilar stationary phase or using another technique such as GC/MS. If the retention times of the peaks on both columns fall within the retention time windows on the respective columns, then the target analyte identification has been confirmed.
  - 7.5.5.1 An additional criterion is applied for the identification and quantitation of PCBs. Identification is based on the characteristic fingerprint

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retention time and shape of the major peaks. Major peaks are defined as those peaks in the Aroclor standard that are at least 25% of the height of the largest Aroclor peak. The sample chromatogram is compared to the individual Aroclor standard chromatograms. Once the Aroclor pattern has been identified, a concentration is then calculated in Target.

- 7.5.5.2 Three to five Aroclor concentrations are calculated using the peak heights of the three to five characteristic peaks of the Aroclor. These three to five concentrations are then averaged to determine the concentration of that Aroclor.
- 7.5.6 When samples are analyzed from a source known to contain specific Aroclors, the results from a single-column analysis may be confirmed on the basis of a clearly recognizable Aroclor pattern.
- 7.5.7 If the response for an analyte exceeds the calibration range of the system, the sample must be diluted and reanalyzed.
- 7.5.8 If peak detection and identification are prevented due to interferences, the hexane extract may need to undergo a cleanup. The extract may be subjected to a sulfur cleanup (method 3660) and/or a sulfuric acid cleanup (method 3665).

**Note:** Samples routinely receive a sulfuric acid clean up. However, for samples from a known site with a clean matrix, a sulfuric acid clean up may not be performed. Whenever a sample receives a cleanup, the associated QC must also be subjected to the same cleanup(s) and reanalyzed.

7.5.9 When a GC system is determined to be out of control because either a CV cannot pass or a six point calibration does not meet the correlation coefficient criteria, instrument maintenance is likely necessary. Routine instrument maintenance may involve changing the septum, replacing the liner, clipping the pre-column, or replacing the column. This information is recorded in the instrument run log (Figure 1). When an instrument requires more severe maintenance like replacing the ECD or an electronic board, this information is written in the instrument maintenance logbook.

#### 7.6 Calculations

7.6.1 The concentration of an analyte is calculated by using the calibrated curve that is prepared in Target. When an analyte is identified, Target displays a concentration after the file is processed through the appropriate calibration method. Aroclor quantitation is accomplished by the method described in

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section 7.5.4.1.1. However, if a sample contains more than one Aroclor, a peak common to both analytes must not be used to quantitate either compound.

7.6.2 The concentrations from the reports are then incorporated with the extraction data to arrive at a final concentration.

Water: Concentration (ug/L) = (C) (Vt)/(Vs)

Soil/Sediment: Concentration (mg/kg) = (C) (Vt)/ (Ws) (D)

where, C = concentration calculated by Target in ug/ml

Vt = Volume of total extract including any instrument dilutions

Vs = Volume of sample extracted Ws = Weight of sample extracted

D = Decimal total solids

#### 7.7 Data Review

#### 7.7.1 Initial Data Review

The initial data review is accomplished by the analyst who ran the samples. This review is of sufficient quality and detail to provide a list of samples that need to be reanalyzed or diluted and reanalyzed. The initial data review is performed on the detailed quantitation reports of the analyzed samples. This data review examines criteria that directly impact whether or not the sample needs to be reanalyzed and/or extracted. These criteria include:

- ◆ QC criteria for method blank, LCS, MS/MSD, and calibration refer to section 8.0.
- Surrogate recovery
- Chromatography: cleanups, manual integration.
- Target compound detection: quantitation, confirmation, false positives.

The requirement of the GC laboratory is that this initial data review be completed no later than the end of the next workday. After the analyst has completed his or her initial data review, the information is then ready to be processed for reporting. Refer to section 7.8.

#### 7.7.2 Surrogate recovery

All recoveries must meet the most recently laboratory established acceptance limits, which are listed on the GC Laboratory Surrogate Acceptance Limit sheet.

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The sample is evaluated for recoveries of the two surrogates. If the recovery of one surrogate is within the acceptance limit, and the second is out, the data is narrated. If the surrogate recoveries are high for both and the sample contains less than the PQL for all target analytes, the data is narrated. If the surrogate recoveries are low and may be attributable to matrix interference or a matrix effect, the data is narrated. If the surrogate recoveries are low and the sample concentration is less than the PQL for all target analytes and there is no apparent matrix effect, reextract the sample.

For method blanks, if the recoveries of both surrogates are low or high, and the blank does not contain any target analytes above the PQL, and the recoveries of both surrogates in the sample(s) are acceptable, the data is narrated. If the recoveries in the blank are low and it does not contain any target analytes above the PQL, and the recoveries in the samples are acceptable but the sample contains one or more target analytes above the PQL, the sample may be reextracted.

For laboratory control samples (LCS), if the only discrepancy in the extraction batch is with the LCS, and the analyte spike recoveries are acceptable, the data is narrated. If the recoveries of both the surrogates and the analyte spikes are low, the samples may need to be reextracted.

For DoD QSM 4.1, use QC acceptance criteria specified by DoD, if available. Otherwise, use in-house control limits. When the surrogate recoveries fall outside of the acceptance criteria, apply Q-flag to all associated analytes.

#### 7.7.3 Chromatography

The chromatography should be examined for the presence of any non-target peaks, which can be used as an indication of whether or not matrix interference might be influencing surrogate recoveries. If the chromatogram indicates interferences, then a cleanup may be needed. See section 7.5.7.

Manual integrations are to be performed when chromatographic conditions preclude the computer algorithm from correctly integrating the peak of concern. In no instance shall a manual integration be performed solely to bring a peak within criteria.

Each peak of concern is examined by the primary analyst to ensure that the peak was integrated properly by the computer algorithm. Should a manual integration be necessary (for instance, due to a split peak, peak tailing, or incomplete resolution of isomeric pairs), an "m" qualifier will automatically be printed on the quantitation report summary. The analyst will date and initial the "m" on the quanitation report summary and assign a code that indicates

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the reason for the manual integration. Refer to Katahdin SOP QA-812 "Manual Integration on GC/MS, GC, HPLC and IC Datasystems" for more information.

#### 7.7.4 Target Compound Detection

GC analysis relies heavily on the experience of the analyst. Sample chromatograms must be evaluated focusing on scientific judgment, knowledge of the column behavior and matrix effects. The chromatogram from channel A is evaluated with that from channel B. If a target analyte is present on both channels and the concentration is within the calibration range, and the quantitation from both chromatograms agrees within  $\pm 40\%$ , the analyte is considered present in the sample. In cases where the RPD is greater than 40% and the analyte is reported, the analyte must be J-flagged and narrated. The higher of the two concentrations is reported unless matrix interference is causing erroneously high results. In this case report the lower result and narrate. In some cases a non-confirming analyte may be reported. In these cases the analyte must be Q-flagged and narrated...

In order to avoid reporting false positives, identified peaks on a chromatogram may need to be undetected electronically in Target. The possible scenarios are: If an analyte is present on one column but its concentration is below the PQL, if an analyte is present on one column but does not confirm on the other channel, if an analyte is present on both columns but the concentrations differ by more than 40%, or if an analyte is present but its retention time is  $\pm 0.04$  minutes or more than the retention time of the analyte in the preceding CV. The GC Analyst must rely on technical experience in reviewing chromatograms in determining if a hit is an actual analyte or a false positive.

If reporting data that has an RPD that is >40%, the data must be flagged with a "J" indicating that the result is an estimated value. Sometimes interference on one column (i.e. sulfur) will prevent a target analyte from detection and it is present on the conformational column. In this scenario, the result would be reported from one column and need to be "Q" flagged to indicate that it was not confirmed on a second column.

#### 7.7.5 Reporting

After the chromatograms have been reviewed and any target analytes have been quantitated using Target, the necessary files are brought into KIMS. Depending on the QC level requested by the client, a Report of Analysis (ROA) and additional reports, such as LCS forms and chronology forms, are generated. The package is assembled to include the necessary forms and raw data. The data package is reviewed by the primary analyst and then forwarded

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to the secondary reviewer. The secondary reviewer validates the data and checks the package for any errors. When completed, the package is sent to the Department Manager for final review. A completed review checklist is provided with each package. The final data package from the Organics department is then processed by the Data Management department.

#### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

Refer to Table 1 and to details in this section for a summary of QC requirements, acceptance criteria, and corrective actions. These criteria are intended to be guidelines for analysts. The criteria does not cover all possible situations. If any of the QC requirements are outside the recovery ranges listed in this section or in Table 1, all associated samples must be evaluated against all the QC. In some cases data may be reported, but may be reanalyzed in other cases. Making new reagents and standards may be necessary if the standardization is suspect. The corrective actions listed in this section and in Table 1 may rely on analyst experience to make sound scientific judgments. These decisions are based on holding time considerations, client and project specific Data Quality Objectives and on review of chromatograms. The supervisor, Operations Manager, and/or Quality Assurance Officer may be consulted to evaluate data. Some samples may not be able to be reanalyzed within hold time. In these cases "qualified" data with narration may be advisable after consultation with the client.

In some cases the standard QC requirements listed in this section and in Table 1 may not be sufficient to meet the Data Quality Objectives of the specific project. Much of the work performed at the lab is analyzed in accordance with specific QC requirements spelled out in a project specific Quality Assurance Project Plan (QAPP) or in a program specific Quality Systems Manual (QSM). The reporting limits, acceptance criteria and/or corrective actions may be different than those specified in this SOP. In these cases the appropriate information will be communicated to the Department Manager and/or senior chemists before initiation of the analyses so that specific product codes can be produced for the project. In addition, the work order notes for each project will describe the specific QAPP or QSM to be followed.

- 8.1 For each analytical batch (up to 20 samples), a method blank, laboratory control sample (LCS), matrix spike and matrix spike duplicate are analyzed. They are carried through all stages of the sample preparation and analysis steps.
  - 8.2 Spike concentrations: The LCS and the MS/MSD are spiked at the same concentration with Aroclor 1660. The spike concentrations are:

Compound	WATER ug/L	SOILS mg/kg
Aroclor 1660	5.0	0.17

The surrogate spike concentrations in the final extract are:

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Compound	WATER ug/ml	SOILS ug/ml
Tetrachloro-m-xylene(TCX)	0.10	0.10
DCB	0.10	0.10

8.3 LCS and MS/MSD acceptance criteria and Corrective Action: All QC samples are calculated for percent recovery of the spiked analyte. The recoveries are compared to laboratory established acceptance limits. The LCS acceptance limits for PCBs are established for both water and soil matrices. The MS/MSD acceptance limits for PCBs use the respective matrix LCS acceptance limits. Separate limits for MS/MSD pairs are not calculated because of the varying matrices involved. In addition many of the MS/MSD data points cannot be used (i.e. recoveries not calculable due to a matrix effect).

If any spike compound in the laboratory control sample falls outside of the established recovery acceptance limit window, the QC sample is considered to be out of control and any sample that is associated should be evaluated with other QC elements to determine the corrective action. If the recovery is high and the associated samples do not contain the specific compound(s), the data can possibly be accepted with narration. In other cases, the associated samples must be extracted.

If a spike compound is outside of the acceptance limits in the matrix spike sample but is acceptable in the LCS, the data is considered acceptable. The cause of the failure is possibly attributable to matrix interference. However, if the compound fails in both the LCS and the MS/MSD, the result for that analyte is suspect and may not be reported for regulatory compliance purposes.

For DoD QSM 4.1, use QC acceptance criteria specified by DoD, if available. Otherwise use in-house control limits. In-house control limits must not be greater than  $\pm$  3 times the standard deviation of the mean LCS recovery. If the LCS fails the acceptance criteria, correct problem, then reprep and reanalyze the LCS and all samples in the associated preparatory batch for failed analytes, if sufficient sample material is available. If reanalysis cannot be performed, data must be qualified and explained in the narrative. Apply Q-flag to specific analyte(s) in all samples in the associated preparatory batch.

For MS, when applying DoD QSM 4.1, apply J-flag to specific analyte(s) also in parent sample, if acceptance criteria not met. RPD must be </= 30% between MS and MSD.

8.4 Surrogate acceptance criteria and Corrective Action: Surrogate recoveries are calculated on all samples, blanks and spikes. The recoveries are compared to laboratory established acceptance limits.

When a sample has a surrogate that falls outside of the laboratory established acceptance limit window, the problem should be investigated. If the recovery looks like it

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is affected by the sample matrix, the sample may be reinjected to confirm matrix interference. When a sample has no detectable surrogate recovery, the sample should be reextracted.

For DoD QSM 4.1, use QC acceptance criteria specified by DoD, if available. Otherwise, use in-house control limits. When the surrogate recoveries fall outside of the acceptance criteria, apply Q-flag to all associated analytes.

8.5 Non-conformance Report: Whenever data is not acceptable because of a failing LCS or surrogate recovery, a non-conformance report (NCR) must be initiated as soon as possible to document resolution.

#### 9.0 METHOD PERFORMANCE

The method detection limit (MDL) is defined as the minimum concentration of a substance that can be measured and reported with 99% confidence that the value is above zero. The MDLs are determined prior to sample analysis per type of instrument and filed with the Organic Department Manager and with the QAO. Refer to the current revision of Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, for procedures on determining the MDL.

Refer to the current revision of Method 8082 for other method performance parameters and requirements.

#### 10.0 APPLICABLE DOCUMENTS/REFERENCES

Test Methods for Evaluating Solid Waste, Physical/Chemical Methods, SW-846, 3rd Edition. Final Update IV, dated February, 2007, Method 8082A.

The National Environmental Laboratory Accreditation Conference (NELAC) Standards, June 2003.

The NELAC Institute, Laboratory Accreditation Standards, Volume 1, Management and Technical Requirements for Laboratories Performing Environmental Analysis, 10/06/2010.

Department of Defense Quality Systems Manual for Environmental Laboratories (DOD QSM), Current Version.

Katahdin Analytical Services, Inc., SOP CA-101, Equipment Maintenance and Troubleshooting, current revision.

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Katahdin Analytical Services, Inc., SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, current revision.

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Table 2	DoD QSM Version 4.1 QC Requirements
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#### TABLE 1

#### QC REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action
6pt calibration of Aroclor 1660, 1242, 1248, 1254 and mid-point cal of other Aroclors	Initial cal prior to sample analysis	Average Model – at least 5 points, % RSD = 20% Linear Model – at least 5 points, correlation coefficient (r) <math \geq 0.990 Quadratic Model – at least 6 pt calibration, coefficient of determination ( $r^2$ ) $\geq$ 0.990	(1) Repeat Initial calibration (2) If single pt cal Aroclor is identified in analysis of sample,5 or 6-pt calibration (depending on calibration model) of identified compound with reanalysis of sample.
Independent Calibration Verification	Immediately following calibration	± 20 % D	(1) Reanalyze standard (2) Reprep standard (3) Reprep standard from fresh stock.
CCV	After every 10 samples; If calibration curve previously analyzed, analyze daily before samples.	± 15 % D	<ol> <li>Evaluate the samples: If the %D &gt;+15% and sample results are <pql, li="" narrate.<=""> <li>If %D &gt;±15% only on one channel, narrate. If %D &gt;±15% for closing CV, and is likely a result of matrix interference, narrate.</li> <li>Otherwise, reanalyze all samples back to last acceptable CV.</li> </pql,></li></ol>
Method blank	One per prep batch	No analyte detected >PQL	<ol> <li>Investigate source of contamination</li> <li>Evaluate the samples and associated QC: i.e. If the blank results are above the PQL, report sample results which are <pql or=""> 10X the blank concentration.</pql></li> <li>Otherwise, reprep a blank and the remaining samples.</li> </ol>
LCS	One per prep batch of twenty or fewer samples	Laboratory statistically derived limits.	<ol> <li>Evaluate the samples and associated QC: i.e. If an MS/MSD was performed and acceptable, narrate.</li> <li>If an LCS/LCSD was performed and only one of the set was unacceptable, narrate.</li> <li>If the surrogate recoveries in the LCS are also low but are acceptable in the blank and samples, narrate.</li> <li>If the LCS recovery is high but the sample results are <pql, li="" narrate.<=""> <li>Otherwise, reprep a blank, QC and the remaining samples.</li> </pql,></li></ol>
Matrix Spike\ Matrix Spike Duplicate	One for every set of 20 samples	Same as for LCS	Evaluate the samples and associated QC: i.e. If the LCS results are acceptable, narrate.     If both the LCS and MS/MSD are unacceptable reprep samples and QC.

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#### TABLE 1 (cont.)

#### QC REQUIREMENTS

QC Check	Minimum	Acceptance Criteria	Corrective Action	
	Frequency			
Sample Duplicate	One sample duplicate per ten samples if requested	RPD <u>&lt;</u> 20	<ul><li>(1) If lab QC in criteria and matrix interference suspected, flag data or narrate</li><li>(2) Otherwise, reanalyze</li></ul>	
Demonstration of analyst proficiency – 4 replicates	Once per analyst initially and annually thereafter	P&A meet method criteria	(1) Repeat P&A study	
MDL study	Refer to KAS SOP QA-806, "Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications", current revision.			

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# TABLE 2 DOD QSM QC REQUIREMENTS

QC Check	Minimum	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Demonstrate acceptable analytical capability	Prior to using any test method and at any time there is a significant change in instrument type, personnel, test method, or sample matrix.	QC acceptance criteria published by DoD, if available; otherwise, method-specified criteria.	Recalculate results; locate and fix problem, then rerun demonstration for those analytes that did not meet criteria.	Not Applicable (NA).	This is a demonstration of analytical ability to generate acceptable precision and bias per the procedure in Appendix C. No analysis shall be allowed by analyst until successful demonstration of capability is complete.
LOD determination and verification LOQ establishment	Refer to current r	evision of SOP QA-806 evision of SOP QA-806			
and verification Retention time (RT) window width calculated for each analyte and surrogate	At method set- up and after major maintenance (e.g., column change).	RT width is ± 3 times standard deviation for each analyte RT from a 72-hour study.	NA.	NA.	
Minimum five- point initial calibration (ICAL) for all analytes	ICAL prior to sample analysis.	6 point calibration of Aroclors 1016, 1242, 1248, 1254 and 1260 - One of the options below: Option 1: RSD for each analyte ≤ 20%; Option 2: linear least squares regression: r ≥ 0.995; Option 3: non-linear regression: coefficient of determination (COD) r2 ≥ 0.99 (6 points shall be used for second order). Mid point calibration of Aroclors 1221 and 1232; if targets are detected, 6-point calibration is performed.	Correct problem then repeat ICAL.	Flagging criteria are not appropriate.	Problem must be corrected. No samples may be run until ICAL has passed. Calibration may not be forced through the origin. Quantitation for multicomponent analytes such as chlordane, toxaphene, and Aroclors must be performed using a 5-point calibration. Results may not be quantitated using a single point.
Retention time window position establishment for each analyte and surrogate	Once per ICAL and at the beginning of the analytical shift.	Position shall be set using the midpoint standard of the ICAL curve when ICAL is performed. On days when ICAL is not performed, the initial CCV is used.	NA.	NA.	

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#### TABLE 2 (cont)

#### DOD QSM QC REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Second source calibration verification (ICV)	Immediately following ICAL.	All project analytes within established retention time windows. All project analytes within ± 20% of expected value from the ICAL.	Correct problem, rerun ICV. If that fails, repeat ICAL.	Flagging criteria are not appropriate.	Problem must be corrected. No samples may be run until calibration has been verified.
calibration verification (CCV)	Prior to sample analysis, after every 10 field samples, and at the end of the analysis sequence.	All project analytes within established retention time windows. All project analytes within ± 20% of expected value from the ICAL.	Correct problem, then rerun calibration verification. If that fails, then repeat ICAL. Reanalyze all samples since the last successful calibration verification.	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply Q-flag to all results for the specific analyte(s) in all samples since the last acceptable calibration verification.	Problem must be corrected. Results may not be reported without a valid CCV. Flagging is only appropriate in cases where the samples cannot be reanalyzed. Retention time windows are updated per the method.
Method blank	One per preparatory batch.	No analytes detected > ½ RL (> RL for common lab contaminants) and > 1/10 the amount measured in any sample or 1/10 the regulatory limit (whichever is greater). Blank result must not otherwise affect sample results.	Correct the problem. Report sample results that are <lod or="">10x the blank concentration. Reprepare and reanalyze the method blank and all associated samples with results &gt; LOD and &lt; 10x the contaminated blank result. Contact Client if samples cannot be reprepped within hold time.</lod>	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply B-flag to all results for the specific analyte(s) in all samples in the associated preparatory batch.	Problem must be corrected. Results may not be reported without a valid method blank. Flagging is only appropriate in cases where the samples cannot be reanalyzed.
Laboratory control sample (LCS) containing all analytes to be reported, including surrogates	One per preparatory batch.	The laboratory shall use laboratory control limits (CLs) or use DoDgenerated LCS-CLs, if available depending on project requirements. Inhouse CLs may not be greater than ± 3 times the standard deviation of the mean LCS recovery. A number of analytes may fall outside the CL but within marginal exceedance limit depending on the total number of analytes in the LCS.	Correct problem, then reprep and reanalyze the LCS and all samples in the associated preparatory batch for failed analytes, if sufficient sample material is available. Refer to Table G-1 for number of marginal exceedences allowed. Contact Client if samples cannot be reprepped within hold time.	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply Q-flag to specific analyte(s) in all samples in the associated preparatory batch.	Problem must be corrected. Results may not be reported without a valid LCS. Flagging is only appropriate in cases where the samples cannot be reanalyzed.

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#### TABLE 2 (cont.)

#### DOD QSM REQUIREMENTS

QC Check	Minimum	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Matrix spike (MS)	One per preparatory batch per matrix if sufficient sample is available.	For matrix evaluation, use laboratory control limits (CLs) or use DoDgenerated LCS-CLs, if available depending on project requirements.	Examine the project- specific DQOs. Contact the client as to additional measures to be taken.	For the specific analyte(s) in the parent sample, apply J-flag if acceptance criteria are not met.	For matrix evaluation only. If MS results are outside the LCS limits, the data shall be evaluated to determine the source of difference and to determine if there is a matrix effect or analytical error.
Matrix Spike duplicate (MSD)	One per preparatory batch per matrix if sufficient sample is available.	MSD: For matrix evaluation, use laboratory LCS CLs or use DoD- generated LCS CLs, if available depending on project requirements. MS/MSD: RPD ≤ 30%.	Examine the project- specific DQOs. Contact the client as to additional measures to be taken.	For the specific analyte(s) in the parent sample, apply J-flag if acceptance criteria are not met.	The data shall be evaluated to determine the source of difference.
Surrogate spike	All field and QC samples.	The laboratory shall use laboratory CLs or use DoD-generated Surrogate CLs, if available depending on project requirements.	For QC and field samples, correct problem then reprep and reanalyze all failed samples for failed surrogates in the associated preparatory batch, if sufficient sample material is available. If obvious chromatographic interference with surrogate is present, reanalysis may not be necessary. Contact Client if samples cannot be reprepped within hold time.	Apply Q-flag to all associated analytes if acceptance criteria are not met.	Alternative surrogates are recommended when there is obvious chromatographic interference.
Confirmation of positive results (second column or second detector)	All positive results must be confirmed (with the exception of Method 8015).	Calibration and QC criteria same as for initial or primary column analysis. Results between primary and second column RPD ≤ 40%.	NA.	Apply J-flag if RPD > 40%. Discuss in the case narrative.	Use project-specific reporting requirements if available; otherwise, use method reporting requirements; otherwise, report the result from the primary column.
Results reported between DL and LOQ	NA.	NA.	NA.	Apply J-flag to all results between DL and LOQ.	

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### TITLE: ANALYSIS OF PCBs AS TOTAL AROCLORS BY GAS CHROMATOGRAPHY/ ELECTRON CAPTURE DETECTOR (GC/ECD): SW-846 METHOD 8082

# TABLE 3 SUMMARY OF METHOD MODIFICATIONS

TOPIC	KATAHDIN SOP CA-329-13	METHOD 8082, current revision
Procedures	7.4.5 If the calculated retention time window results in a value of 0.03 minutes or less, the laboratory will apply nominal windows. This is done in order to avoid any false negative hits because of the window being to narrow. The windows are: ± 0.07 for all target analytes. By utilizing these windows, a false positive hit may be initially indicated, but an experienced analyst could determine a false positive from scrutinizing the chromatograms. Please note that the use of nominal retention time windows may not be allowable for certain states, federal programs, or clients. South Carolina does not allow the use of nominal limits for compliance work originating in their state. In these cases, a window of ± 0.03 minutes must be used if the established retention time window is less than 0.03 minutes.	9.3 refers to method 8000B section 7.6.3: If the standard deviation of the retention times for a target compound is 0.000 (i.e., no difference between the absolute retention times), then the laboratory may either collect data from additional injections of standards or use a default standard deviation of 0.01 minutes. (Recording retention times to three decimal places rather than only two should minimize the instances in which the standard deviation is calculated as 0.000).
Apparatus/Materials	time window is less than 0.03 minutes.	
Reagents		
Sample Preservation and handling		
QC – Spikes		
QC - LCS		
QC – Accuracy/ Precision		
QC - MDL	PQL Practical Quantitation Level – three to ten times the MDL.	EQL Estimated Quantitation Level – five to ten times the MDL

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# FIGURE 1 EXAMPLE OF INSTRUMENT RUN LOG

GC Labo	oraton	y Instrument	Runlog		Standard		Standard ID	
Instrumer	nt: GC	07		Po	Llubo 1-0	18691	16813	
Amount I	njecte	d: 2 uL	1	A	21M21.0	Plet	ls.	
Column N	Numbe	ers:390	321	A	NN48 1.0	Plezo	lo	
		P		A	C1254 (0	Plete	12	
Method:		SW846 80	82 EPA 608	1	42/600 0.25	168	12	
(circle)				+			T	
Date	Init.	Result File	Sample ID Waryld Tree	Y/N	Analytical Workgroup	Method	Comments	
2/11/13	Ch	746136	PLINE	N	MAG IN COMMON	Palost	where chock	
1	1	1 139	ALLIGOTO	1		1	1	
7	1	J 140	ALINA 1.0	7		-1	1	
Mulis	Ch	746/41	PHINA	N		906057		
1	1	, ///	ALIGGO 10	4	W4/20137		1764, 16, 40 CB ATT	
		143	AL1242 1.0	4	1			
		144	RRING 10	4				
		145	AP1254 1.0	1				
		146	W4120093-1 3500	1			Chrospymas 8.13	
		147	1 -2				81.10 COMOPASSIVALI	
		11/8	Suoquy-1				Jucces Inacting	
		149	7.5 7	1			7 7	
		150	KR-1660 0-15	4			1754416 A	
		151	10/12/10					
		152	O. I SMYLDA	1				
	4	153	Mercy 1.0	1	1	1		
Note	Ob	746 154	PHMA	N		106057		
1	-	155	AFLUVOIA	N	W4120196-1.2	1	ATURA	
-	1	156	ALMIN 1.0	4	1			
	-	152	ALME LO	14	-			
4	1	1 15%	ARRENLO	14		1		
-	+	159	MAISCOSZ-1 3510	+		-		
-	+	160	-2	+		-		
	-	161	V -3	+			tr as	
-	+	162	540792-1	+	-	+	TrueA	
-	-	163	1 -2	+	<del>                                     </del>	-	A927	
-	-	164	-3 -y	+		1	ALUTA	
	-	165	54084-1	+			A JOTA	
1	4	100		+	1-4-	1-9	ATUR	
7	1	1 163	1-2	1		7	TOM	

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#### FIGURE 2

#### DATA REVIEW CHECKLIST

#### PRIMARY REVIEW CHECKLIST

nt:		Primary		Secondary
nod:		Date:	Date:	
S No:	Level:	Initials:	Initials:	
No:		•	Approved	: 🔲
OODOSM (4.1)	DOD W	// LAB. LIMITS 🗆	OUAPP	LAB 🗆
. , ,	RT <u>ND's</u> t			
List all curves that ar	e scanned (	hard copy not included ).	1	
Narrote which OC li	mite wara u	sed for ( Surr., LCS's MS	/MSD's )	П
All needed forms are		sed for ( suit., Des s Ms	(MSD 8.)	
		SDG name (all forms).		
		g (all forms). (Truncated	∪ ).	
Correct file numbers	(all forms).			
Analysis Date Correc	t.			
Extraction Method &	Analysis M	ethod Correct.		
Product list compared	l to ROAs (	compounds & PQLs).		
Chromatogram review	wed for unla	beled peaks (check produ	act list).	
Flagging of all ROAs	correct (F	lorida 🗆 ) ( Florida 🗆 )		
All tunes included (le	vel IV).			
All log book pages in	cluded (Soi	weights, TCLP & SPLP)		
Verify DOD QSM cr	iteria and/o	r Project specific requirer	ments.	
Narrate any method	deviations.	( Blanks, LCS's etc. )		
Sign & Date Manual	integration	( Narrate as needed ).		
Sample I.D's Truncat			Please list KAS #	

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# FIGURE 3 PQLs FOR METHOD 8082

ANAL	Practical Quantitation	Practical Quantitation
YTE	Level (PQL)	Level (PQL)
116	(ug/L)	(ug/kg)
PCB-1016	0.50	17
PCB-1221	0.50	17
PCB-1232	0.50	17
PCB-1242	0.50	17
PCB-1248	0.50	17
PCB-1254	0.50	17
PCB-1260	0.50	17
PCB-1262	0.50	17
PCB-1268	0.50	17

### KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

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		STION OF SOLID SAMPLES BY USEPA ME BY ICP-AES AND GFAA	ETHOD 3	050 FOR METALS
Prepared By	y: .	George Brewer	Date:_	3/98
Approved B	y:			
Group Supe	ervisor:	Swage Brewer	Date:_	01/24/01
Operations	Manager: <sub>_</sub>	Il C. Burton	Date:_	1/24/01
QA Officer:	-	Dutorah J. Nadeau	Date:_	1.24.01
General Ma	nager: <sub>-</sub>	Durant. Lufan	Date:_	1/25/01

#### Revision History:

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
01 3050b	Format changes, added pollution prevention, added MSD, added Spiking instruction—tables	8n	12401	1/24/01
02 3050B	Removed all references/procedures de- voted to GFAA. Added use of digestates for ICPMS analysis. Revised steindard solution names 4 concs. in Tables 34 4 to reflect current practice.	Dn	8:29-02	8.39.02
03 3050B	New Title to include 1 Lm05, 3. Use of digestion blockand polyethylene digestion tubes added to sections 4.0, 7.0 and Table 1. PBS changed from 1.09 water to 1.09 booling chips. Hz02 addition from 3,000 then 7.000 to 5.000, 2.000 then 7.000 figures and Tables updated to reflect correct p	LAD	03/08	03/08
04	Updated Tables 3 and 4 with current 'Spike concentrations and volumes added. Updated Logbook page. Added CA-108 reference for Subsempling information.	LAD	08109	08/08
05	updated Tables 3 and 4 to reflect corrent spiking procedures.	LAY	09/10	04/10

### KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-605-05 Date Issued: 09/10

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TITLE:	ACID DIGESTION OF SOLID SAMP ANALYSIS BY ICP-AES, ICP-MS	LES BY USEPA METHOD 3050 FOR METALS
	acknowledge receipt of this standard ope provided. Return the bottom half of this s	erating procedure by signing and dating both of the heet to the QA Department.
	SAMPLES BY USEPA METHOD 3050 F	SOP CA-605-05, titled ACID DIGESTION OF FOR METALS ANALYSIS BY ICP-AES, ICP-MS
Recipient	ıt:	Date:
	DIN ANALYTICAL SERVICES, INC. ARD OPERATING PROCEDURE	
	SAMPLES BY USEPA METHOD 3050 F	SOP CA-605-05, titled ACID DIGESTION OF OR METALS ANALYSIS BY ICP-AES, ICP-MS
Recipient	nt·	Date <sup>.</sup>

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TITLE: ACID DIGESTION OF SOLID SAMPLES BY USEPA METHOD 3050 FOR METALS ANALYSIS BY ICP-AES, ICP-MS

#### 1.0 SCOPE AND APPLICATION

The purpose of this SOP is to describe the Katahdin Analytical Services, Inc. procedure utilized to dissolve solid matrices and solubilize metals from solid samples prior to analysis for metals by ICP-AES and ICP-MS. This SOP applies to samples prepared by EPA Method 3050, with method modifications as summarized in Table 2.

This procedure applies to all solid sample (e.g. sediments, sludges, soils, and ashes) preparations for ICP-AES and ICP-MS analyses. This method is not a <u>total</u> digestion technique for most samples. It is a very strong acid digestion that will dissolve almost all elements that could become "environmentally available". By design, elements bound in silicate structures are not normally dissolved by this procedure as they are not usually mobile in the environment.

#### 1.1 Definitions

<u>ICP-AES</u> – Inductively Coupled Plasma Atomic Emission Spectroscopy.

<u>ICP-MS</u> – Inductively Coupled Plasma Mass Spectrometry.

<u>LCSS</u> – Laboratory Control Sample for Solids – A standard or solid reference material that has been brought through the sample preparation process.

<u>Matrix Spike</u> – An aliquot of a sample to which a known amount of analyte has been added before digestion.

<u>PBS</u> – Preparation Blank for Solids – An aliquot of reagent water that has been brought through the sample preparation process.

#### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of analysts experienced in the acid digestion of solid samples by USEPA Method 3050 for metals analysis. Each analyst must demonstrate the ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, current revision, "Personnel Training & Documentation of Training".

It is the responsibility of all Katahdin technical personnel involved in the acid digestion of solid samples by USEPA Method 3050 to read and understand this SOP, to adhere to the procedures outlined, and to properly document their data in the appropriate lab notebook. Any deviations from the procedure or irregularities with the samples should also be recorded in the lab notebook and reported to the responsible Department Manager or designated qualified data reviewer.

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### TITLE: ACID DIGESTION OF SOLID SAMPLES BY USEPA METHOD 3050 FOR METALS ANALYSIS BY ICP-AES, ICP-MS

It is the responsibility of the Department Manager to ensure that technical personnel perform acid digestions in accordance with this SOP and to confirm that their work is properly documented through periodic review of the associated logbooks.

#### 1.3 Safety

The acids used in this procedure are highly corrosive and reactive, and spiking standards contain toxic metals. The toxicity and reactivity of client samples are usually unknown, so samples should always be assumed to present a contact hazard. To reduce or eliminate exposure to potentially harmful chemicals, lab coats, gloves, and safety glasses or goggles must be worn whenever handling samples or reagents. Additional safety apparel, including face shields, aprons, dust masks, and shoe protectors, is available in the Metals prep lab and should be worn whenever circumstances warrant.

Acids should be added to samples slowly and carefully, while watching for reactions. This should be done under a hood, in case harmful fumes are evolved.

Hood sashes should be lowered as far as possible whenever beakers are being heated on a hot plate. Use caution when handling hot beakers.

Personnel are required to read the Katahdin Hazrdous Waste Management Plan and Safety Manual before performing this procedure, and must be familiar with the general rules for laboratory safety, personal hygiene, housekeeping, and use of protective clothing and equipment.

#### 1.4 Pollution Prevention/Waste Disposal

Whenever possible, laboratory personnel should use pollution prevention techniques to address their waste generation. Refer to the current revision of the Katahdin Hazardous Waste Management Program for further details on pollution prevention techniques.

Excess spiking solutions must be emptied into the corrosive waste carboy located in the Metals prep lab for subsequent appropriate disposal in accordance with the Katahdin Hazardous Waste Management Plan and Safety Manual.

Sample digestates should be stored for a minimum of 60 days after digestion to allow for analysis, and reanalysis if necessary. Digestates older than 60 days may be emptied into the corrosive waste carboy in the Metals prep lab for subsequent appropriate disposal in accordance with the Katahdin Hazardous Waste Mnagement Plan and Safety Manual.

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#### 2.0 SUMMARY OF METHOD

A representative 1 to 2 g (wet weight) sample is digested with repeated additions of nitric acid and hydrogen peroxide. Hydrochloric acid is added to the initial digestate and the sample is refluxed. The digestate is then filtered and diluted to a final volume of 100 mL.

#### 3.0 INTERFERENCES

Interferences are discussed in the applicable analytical SOPs.

#### 4.0 APPARATUS AND MATERIALS

- 4.1 Digestion vessels. If digestion is performed using a hot plate, the appropriate digestion vessels are 100 mL pre-cleaned Griffin beakers (cleaned according to the current revision of SOP CA-100, "Labware Cleaning" and CA-602, "Glassware Preparation and Sample Preservation for Trace Element Analyses"). If digestion is performed using a block digester, the appropriate digestion vessels are new 70 mL disposable graduated polyethylene digestion tubes with attached snap lids.
- 4.2 Ribbed watch glasses. If digestion is performed using a hot plate, 75 mm diameter glass watch glasses (pre-cleaned as above) are used. If digestion is performed using a block digester, 40 mm diameter disposable polyethylene watch glasses are used.
- 4.3 Adjustable volume automatic pipets covering the range from 10 uL to 1000 uL and disposable pipet tips; calibrated Finn pipets or Eppendorf pipets are acceptable.
- 4.4 Disposable graduated polystyrene specimen containers with pouring lips, 200 mL capacity.
- 4.5 Hot plate or block digester, griddle, or other heating source adjustable and capable of maintaining a temperature of 95°C ± 5°C. Heating sources must be numbered for easy identification.
- 4.6 Device for measuring hot plate temperature, consisting of a flask or digestion vessel in which the bulb of a thermometer is immersed in sand or water. The temperature of each hot plate used is measured and recordedeach day. The hot plate identification number and the measured temperature are recorded on the sample preparation logbook sheet.
- 4.7 Plastic funnels, pre-cleaned as in Section 4.1.

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- 4.8 Filter funnel holders, capable of suspending plastic funnels above disposable specimen containers.
- 4.9 Polyethylene wash bottles for dispensing reagent water and 5% HNO<sub>3</sub>.
- 4.10 Filter paper, Whatman No. 41 or equivalent. Filters are acid-washed immediately prior to use as follows. Place a pre-cleaned funnel in the funnel holder and put a disposable plastic specimen container under the funnel to collect the rinsates. Place a folded filter in the funnel and rinse three times with approximate 10 mL volumes of 5% HNO<sub>3</sub>, making sure the entire surface of the filter is wetted each time and allowing each rinse to drain completely before continuing. Then rinse three times with approximate 25 mL volumes of reagent water, again allowing each rinse to drain completely. Discard the rinsates into the appropriate waste container. The acid-washed filter is now ready for use.
- 4.11 Polyethylene sample containers with screw caps or graduated polyethylene sample containers with attached snap lids, 125 mL capacity.
- 4.12 Repipetters (adjustable repeating pipetters with reservoirs) for dispensing concentrated nitric acid, 1:1 HNO<sub>3</sub>, and concentrated HCl.
- 4.13 Analytical balance capable of reading to 0.01 gram.
- 4.14 Spatulas, scoops, or spoons; plastic or stainless steel, rinsed with 5% HNO<sub>3</sub> and reagent water. Disposable tongue depressors may be used and do not require to be rinsed.

#### 5.0 REAGENTS

- 5.1 Concentrated nitric acid, HNO<sub>3</sub> trace metals grade.
- 5.2 Concentrated hydrochloric acid, HCl trace metals grade.
- 5.3 Reagent water water that meets the performance specifications of ASTM Type II water (ASTM D1193).
- 5.4 Nitric acid, 1:1. Add a volume of concentrated HNO<sub>3</sub> to an equivalent volume of reagent water and swirl gently to mix.
- 5.5 Nitric acid, 5% v/v. Add 25 mL concentrated HNO<sub>3</sub> to 475 mL reagent water in a 500 mL wash bottle. Cap, point the dispensing tip into a sink, and shake gently to mix.
- 5.6 30% hydrogen peroxide  $(H_2O_2)$  spectrometric grade.
- 5.7 Multielement spiking solutions (see Table 3 for a list of required spiking solutions).

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5.8 Solid reference material – a soil containing all the elements of interest, with empirically established method-specific recoveries and acceptance limits for all analytes. Solid reference materials are purchased with documentation of analysis provided by the vendor. See Figure 4 for an example certificate of analysis for a solid reference material.

#### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

Samples should be collected in clean plastic or glass containers. Samples must be refrigerated (4°C ±2°C) upon receipt by the laboratory. The holding time for solid samples is 6 months from the date of sample collection.

#### 7.0 PROCEDURE

The procedure described below is condensed for quick reference in Table 3.

#### SAMPLE PREPARATION

- 7.1 Prior to performing the digestion, make a list of the samples that are to be digested. Enter digestion information (Katahdin Sample Numbers, QC Batch ID, preparation date, analyst initials, etc.) into the ACCESS computer spreadsheet. Print out a copy of the spreadsheet (see Figure 2 for an example). Hand label the digestate vessels
- 7.2 If using glass beakers as the digestion vessels, submerge previously cleaned beakers and watch glasses three times into a 10% nitric acid bath, then rinse three times with reagent water. The polyethylene digestion tubes used in conjunction with the block digeter do not require acid rinsing or precleaning. Label the digestion vessels with sample numbers.
- 7.3 Weigh 1 to 2 g of well-mixed sample into a properly cleaned, labeled, and tared Griffin beaker or polyethylene digestion tube. Record (hand write) the weight of each sample on the printout of the digestion spreadsheet. Refer to Katahdin Analytical Services SOP CA-108, current revision "Basic Laboratory Technique" for more information on subsampling.
- 7.4 Weigh an appropriate amount of solid reference material to a clean, labeled, and tared Griffin beaker or polyethylene digestion tube to serve as a laboratory control sample.
- 7.5 Add spike solutions to matrix spike samples (refer to Tables 3 and 4 for spiking instructions).

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- 7.6 Using repipetters, add 10 mL of 1:1 HNO<sub>3</sub>, mix the slurry. Cover with a ribbed watch glass and place on heat source. Gently heat the sample to 95°C ± 5 °C and reflux for 10 to 15 minutes without boiling. Remove the digestion vessel from the heat source and cool the sample.
- 7.7 Add 5 mL of concentrated HNO<sub>3</sub> to the sample, replace the watch glass, and reflux for 30 minutes. If brown fumes are generated, indicating oxidation of the sample by HNO<sub>3</sub>, repeat this step (addition of 5 mL of concentrated HNO<sub>3</sub>) until no brown fumes are given off by the sample, indicating complete reaction by HNO<sub>3</sub>.
- 7.8 Continue heating the sample at 95°C ± 5°C without boiling until the digestate has evaporated to approximately 5 to 10 mL or until two hours have elapsed, whichever occurs first. Do not allow the sample to go to dryness. Remove the digestion vessel from the heat source and cool the sample.
- 7.9 Add 2 mL of reagent water and 2 mL of 30% H<sub>2</sub>O<sub>2</sub> to the sample, replace the watch glass, and heat gently on the heat source to start the peroxide reaction. Continue heating until effervescence subsides.
- 7.10 Add an additional 2 mL of 30% H<sub>2</sub>O<sub>2</sub> to the sample, replace the watch glass, and heat gently on the heat source to start the peroxide reaction. Continue heating until effervescence subsides.
- 7.11 Add an additional 6 mL of 30%  $H_2O_2$  in 1-mL aliquots with warming until the effervescence is minimal or until the general sample appearance is unchanged.
- 7.12 Continue heating the sample at 95°C ± 5°C without boiling until the digestate has evaporated to approximately 5 to 10 mL or until two hours have elapsed, whichever occurs first. Do not allow the sample to go to dryness. Remove the sample from the heat source and cool.
- 7.13 Add 10 mL of concentrated HCl to the digest from 7.12, replace the watch glass, and reflux at  $95^{\circ}$ C  $\pm$   $5^{\circ}$ C for 15 minutes. Remove the sample from the heat source and cool.
- 7.14 Use a pre-cleaned funnel and acid-rinsed filter paper to filter the digestate into a clean graduated polystyrene specimen container or graduated polyethylene sample container with attached snap lid. Using a wash bottle, rinse the digestion vessel with reagent water and add the rinsates to the filter apparatus. After all of the liquid in the filter has drained into the specimen container, thoroughly rinse the filter three times with small (5-10 mL) volumes of reagent water, allowing the liquid to drain completely after each rinse. Using the graduations on the specimen container or snap-lid container, dilute to 100 mL with reagent water. If a specimen container has been used, transfer the contents to the corresponding labeled polyethylene sample bottle, cap the bottle, and discard the empty specimen container. If a snap-lid

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container has been used, close and secure the snap-lid. Shake the container gently to mix. The digestate is now ready for ICP-AES or ICP-MS analysis.

- 7.15 Review the ACCESS computer spreadsheet for accuracy. If any information is incorrect, make the necessary changes to the computer spreadsheet and print out a corrected copy. Do not discard the original copy of the spreadsheet. Record (hand write) reagent lot numbers, spiking information, and heat source temperature in the appropriate spaces on the spreadsheet. Record any method deviations, irregularities with the samples, or other pertinent observations at the bottom of the page, and sign and date the spreadsheet. Bind all copies of the spreadsheet in the sample preparation log. An example sample preparation logbook page (ACCESS spreadsheet) is included as Figure 2.
- 7.15 Reopen the electronic ACCESS spreadsheet for the digestion and transcribe the sample weights from the handwritten, bound copy into the electronic copy. The information in this electronic spreadsheet will later be imported into the ACCESS metals database and used to calculate sample concentrations on a weight basis.
- 7.16 Place each batch of digestates in a box labeled with the QC Batch ID, and put the box of digestates in the metals digestates storage area.

#### **CALCULATIONS**

7.17 Analytical results for solid samples are reported on a dry weight basis. Total solids are determined by the Wet Chemistry Group, and are recorded in spreadsheets that are electronically imported into the Access metals database. Final dry weight concentrations are calculated by the Access database as follows:

Concentration (mg/kg dry weight) =  $(C \times V) / (W \times S)$ 

where: C = Measured concentration (mg/L)

V = Digestate final volume (L)W = Sample wet weight (kg)

S = % Solids/100

#### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

USEPA Method 3050 requires the laboratory to perform specific quality control checks to assess laboratory performance and data quality. Minimum frequencies, acceptance criteria, and corrective actions for these control checks are tabulated in Table 1 and are described below. Table 1 criteria are intended to be guidelines for analysts. The table does not cover all possible situations. If any of the QC requirements are outside the recovery ranges listed in Table 1, all associated samples must be evaluated against all the QC. In some cases data may be reported, but may be reanalyzed in other cases. Making new reagents and

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standards may be necessary if the standardization is suspect. The corrective actions listed in Table 1 may rely on analyst experience to make sound scientific judgements. These decisions are based on holding time considerations and client and project specific Data Quality Objectives. The Department Manager, Operations Manager, and/or Quality Assurance Officer may be consulted to evaluate data. Some samples may not be able to be reanalyzed within hold time. In these cases "qualified" data with narration may be advisable after consultation with the client.

- 8.1 At least one preparation blank for soils (PBS) is processed concurrently with each digestion batch of 20 or fewer samples, and is used to assess contamination resulting from the digestion procedure. The PBS consists of a 1.0 g of boiling stones that is digested using the same reagents as those used to digest associated samples. Refer to the appropriate analytical SOP for PBS acceptance criteria and corrective actions.
- 8.2 At least one laboratory control sample for soils (LCSS) is processed concurrently with each digestion batch of 20 or fewer samples. The LCSS consists of an aliquot of a solid reference material for which the concentrations of the analytes of interest have been empirically established (solid-matrix LCSS), or an aliquot of reagent water that is spiked to contain all analytes of interest at known concentrations (aqueous-matrix LCSS). The solid reference material should normally be used as the LCSS, unless a particular client or analytical program requires that spiked reagent water be used. The LCSS is digested using the same reagents as those used to digest associated samples. Directions for spiking the aqueous-matrix LCSS are contained in Table 3. The measured analyte recoveries for the LCSS are used to assess digestion method performance. Refer to the appropriate analytical SOP for LCSS recovery acceptance criteria and corrective actions.
- 8.3 Matrix spike samples are processed along with each digestion batch at a minimum frequency of one per digestion batch. A matrix spike sample consists of an aliquot of a sample that is fortified with known amounts of all analytes of interest prior to digestion. Matrix spike recoveries are used to assess the biasing effects of sample matrix on digestion and analysis performance. Directions for spiking matrix spike samples are contained in Figure 2. Refer to the appropriate analytical SOP for matrix spike recovery acceptance criteria and corrective actions.
- 8.4 Matrix spiked duplicate samples are processed concurrently with each digestion batch at a minimum frequency of one per digestion batch. Matrix spiked duplicate samples are used to assess the precision of the digestion and analysis methods. Refer to the appropriate analytical SOP for matrix spike duplicate precision acceptance criteria and corrective actions.

<u>NOTE</u>: Clients may choose specific samples for matrix spike and duplicate analysis; otherwise, the choice is left to the person performing the digestion.

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8.5 The quality control measures and frequencies described above are minimum requirements. Individual clients and analytical programs may impose additional QC requirements.

#### 9.0 METHOD PERFORMANCE

Refer to the applicable instrumental analysis SOP for method performance information.

#### 10.0 APPLICABLE DOCUMENTS/REFERENCES

"Test Methods for the Evaluation of Solid Waste," United States Environmental Protection Agency, SW-846, Third Edition, Final Update III, 12/96, Method 3050B.

Department of Defense Quality Systems Manual for Environmental Laboratories (DOD QSM), Version 4.1, 04/22/09.

The National Environmental Laboratory Accreditation Conference (NELAC) Standards, June 2003.

#### LIST OF TABLES AND FIGURES

Table 1	QC Requirements – Method 3050
Table 2	Summary of Method Modifications – Method 3050
Table 3	Preparation of Matrix Spikes and Spiking Solutions
Table 4	Element Concentrations in ICP-AES Matrix Spikes and Their Component Spiking
	Solutions
Figure 1	Procedure Condensation – Method 3050
Figure 2	Example Page from Metals Sample Preparation Logbook
Figure 3	Example Certificate of Analysis for Solid Reference Material

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# TABLE 1 QC REQUIREMENTS – METHOD 3050

Method	QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action
3050	Preparation Blank for Solids (PBS)	One per prep batch of 20 or fewer samples.	Refer to analytical method.	Refer to analytical method.
	Laboratory Control Sample for Solids (LCSS)	One per prep batch of 20 or fewer samples.	Refer to analytical method.	Refer to analytical method.
	Matrix Spike Sample	One per prep batch.	Refer to analytical method.	Refer to analytical method.
	Matrix Spike Duplicate Sample	One per prep batch.	Refer to analytical method.	Refer to analytical method.
	Demonstration of analyst proficiency	One-time demonstration by each analyst performing the method.	Must pass all applicable QC for method.	Repeat analysis until able to perform passing QC; document successful performance in personal training file.

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# TABLE 2 SUMMARY OF METHOD MODIFICATIONS – METHOD 3050

Topic	Katahdin SOP CA-605-02	Method 3050, current revision
Apparatus /Materials	Digestion performed in 100 mL Griffin beaker or 70 mL polyethylene tube.     Graduated disposable plastic cup or 120 mL polyethylene tube used to bring digestate to final volume.	Digestion performed in 250 mL Griffin beaker.     Volumetric flask used to bring digestate to final volume.
Procedure	<ol> <li>Digestate volume reduced to 5 to 10 mL prior to filtering.</li> <li>After filtration, the filters are rinsed three times with reagent water.</li> <li>30% H<sub>2</sub>O<sub>2</sub> is added in two 2 mL aliquots and then six 1 mL aliquots.</li> </ol>	Digestate volume reduced to 5 mL prior to filtering.     After filtration, the filters are rinsed twice with reagent water.     30% H <sub>2</sub> O <sub>2</sub> is added in one 3 mL aliquot and then seven 1 mL aliquots.

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TABLE 3

PREPARATION OF MATRIX SPIKES AND SPIKING SOLUTIONS FOR DIGESTION OF SOLID SAMPLES BY USEPA METHOD 3050

Sample or Solution Name	Component Solution Name	Source of Component	Amount of Component Added per 100 mL Final Volume (mL)
	CLPP-SPK-1	Inorganic Ventures(IV)	0.10
Matrix Child for ICD ACC	CLPP-SPK-INT1	Lab Prepared (see below)	1.00
Matrix Spike for ICP-AES	CLPP-SPK-INT2	Lab Prepared (see below)	1.00
	1000 mg/L Uranium Std.	IV or High Purity Standards	0.01

Sample or Solution Name	Component Solution Name	Source of Component	Amount of Component Added per 100 mL Final Volume (mL)
	1000 mg/L As,Pb,Sb,Se,Tl	High Purity Standards	1.0 each
	1000 mg/L Cd	High Purity Standards	2.5
CLPP-SPK-INT1	10000 mg/L K	High Purity Standards	10.0
	10000 mg/L Na	High Purity Standards	7.5
	10000 mg/L Mg	High Purity Standards	5.0
	10000 mg/L Ca	High Purity Standards	2.5
	1000 mg/L Mo	IV or High Purity Standards	3.0
CLPP-SPK-INT2	1000 mg/L B,Li,Sn,Sr,Ti	IV or High Purity Standards	5.0 each
	10000 mg/L Si	High Purity Standards	5.0

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TABLE 4

ELEMENT CONCENTRATIONS IN ICP-AES MATRIX SPIKES AND THEIR COMPONENT SPIKING SOLUTIONS FOR DIGESTION OF SOLID SAMPLES BY METHOD 3050

	CONCENTRATION IN SOLUTION, mg/L					
	Matrix	CLPP-	CLPP-	CLPP-	CLPP-	1000 mg/L
Element	Spike	SPK-1	SPK-4	SPK-INT1	SPK-INT2	Ŭ
Aluminum	2.000	2000				
Antimony	0.100		100	10		
Arsenic	0.100		4	10		
Barium	2.000	2000				
Beryllium	0.050	50				
Boron	0.500				50	
Cadmium	0.250		5	25		
Calcium	2.500			250		
Chromium	0.200	200				
Cobalt	0.500	500				
Copper	0.250	250				
Iron	1.000	1000				
Lead	0.100		2	10		
Lithium	0.500				50	
Magnesium	5.000			500		
Manganese	0.500	500				
Molybdenum	0.300				30	
Nickel	0.500	500				
Potassium	10.000			1000		
Selenium	0.100		5	10		
Silicon	5.000				500	
Silver	0.050	50				
Sodium	7.500			750		
Strontium	0.500				50	
Thallium	0.100		5	10		
Tin	0.500				50	
Titanium	0.500				50	
Uranium	0.100				† †	1000
Vanadium	0.500	500			†	
Zinc	0.500	500				

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#### FIGURE 1

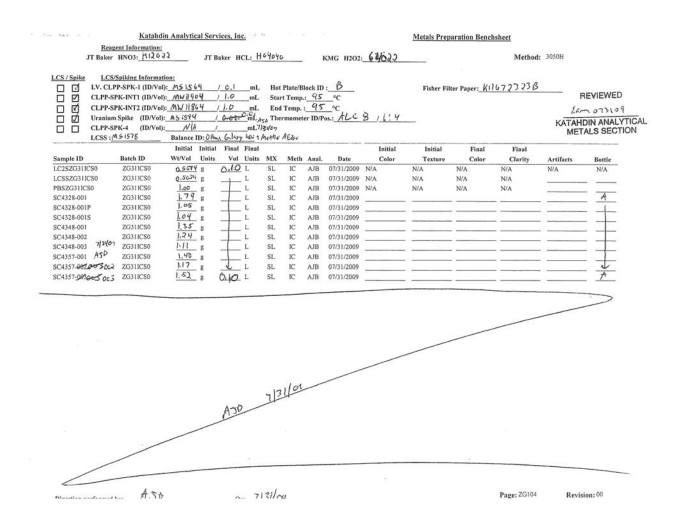
#### PROCEDURE CONDENSATION - METHOD 3050

- 1. Prepare and print out ACCESS spreadsheet.
- 2. If performing digestion on a hot plate, rinse 250 mL Griffin beakers and ribbed watch glasses 3 times in acid bath. Then rinse beakers and watch glasses 3 times with laboratory reagent grade water. If perfoming digestion with block digester, polyethylene digestion tubes do not require precleaning.
- 3. Label digestion vessels (beakers or polyethylene sample tubes) with sample numbers.
- 4. Weigh 1 to 2 g of well-mixed sample into tared digestion vessels. Record sample weights.
- 5. Add spike solutions to matrix spike samples.
- 6. Add 10 mL 1:1 HNO<sub>3</sub> to samples and cover with watch glasses.
- 7. Reflux for 10 to 15 minutes at  $95^{\circ} \pm 5^{\circ}$  C. without boiling. Cool samples.
- 8. Add 5 mL conc. HNO3, cover beakers, and reflux for 30 minutes.
- 9. Repeat Step 8 as necessary until digestion is complete.
- 10. Reduce sample volumes to 5 to 10 mL or heat for 2 hours, whichever occurs first.
- 11. Cool sample and add 2 mL reagent water and 2 mL 30% H<sub>2</sub>O<sub>2</sub>. Heat gently until effervescence subsides.
- 12. Cool sample and add 2 mL 30% H<sub>2</sub>O<sub>2</sub>. Heat gently until effervescence subsides.
- 13. Cool samples and add 6 mL of 30% H<sub>2</sub>O<sub>2 in 1 mL aliquots.</sub> Heat gently until effervescence subsides.
- 14. Reduce sample volumes to 5 to 10 mL or heat for 2 hours, whichever occurs first.
- 15. Add 10 mL conc. HCl and reflux for 10 to 15 minutes at  $95^{\circ} \pm 5^{\circ}$  C.
- 16. Cool sample and filter into graduated specimen container. Bring to volume with reagent water and transfer to labeled polyethylene bottle.
- 17. Enter sample weights into ACCESS spreadsheet.

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# FIGURE 2 EXAMPLE PAGE FROM METALS SAMPLE PREPARATION LOGBOOK



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#### FIGURE 3

#### EXAMPLE CERTIFICATE OF ANALYSIS FOR SOLID REFERENCE MATERIAL



M51475

#### DataPacK™

Lot No. D051-540

#### Trace Metals in Soil

Catalog No. 540

#### Certification

Total	Certified	Performance
Concentration 1	Value 2	Acceptance Limits <sup>™</sup> 3
(mg/Kg)	(mg/Kg)	(mg/Kg)
	(9,9)	(mg/kg)
55600*	7870	4630 - 11100
160		D.L 149
		234 - 344
		174 - 247
		45.2 - 63.6
		58.8 - 124
		82.9 - 119
		2970 - 4390
		180 - 268
		82.7 - 119
		73.3 - 103
24400*		6610 - 24900
184		129 - 187
3780*		1760 - 2750
703		343 - 497
5.32		3.42 - 6.87
		55.5 - 83.7
137		99.1 - 141
33000*		2200 - 3800
		101 - 159
127		68.9 - 139
15600*		692 - 1470
		90.5 - 135
		72.8 - 115
		104 - 194
		116 - 453
		85.1 - 137
		215 - 329
	Concentration 1 (mg/Kg)  55600* 150 316 869 60.9 129 114 9750* 249 113 94.9 24400* 184 3780* 703 5.32 80.2 137 33000* 146	Concentration 1 (mg/Kg) (mg/Kg)  55600* 7870 160 70.5 316 289 869 211 60.9 54.4 129 91.3 114 101 9750* 3680 249 224 113 101 94.9 88.0 24400* 15700 184 158 3780* 2260 703 420 5.32 5.18 80.2 69.6 137 120 33000* 3000 146 130 127 104 15600* 1080 326 113 106 94.0 175 149 3100* 284

	Total	Certified	Performance
Method 3050 HNO3, H2O2	Concentration 1	Value 2	Acceptance Limits™ 3
	mg/Kg	mg/Kg	mg/Kg
Parameter		9,9	mg/kg
aluminum	55600*	7380	4440 - 10300
antimony	160	75.2	
arsenic	316	284	D.L 198 225 - 343
barium	869	217	177 - 257
beryllium	60.9	53.6	42.7 - 64.5
boron	129	89.5	
cadmlum	114	103	58.9 - 120
calcium	9750*	3540	83.6 - 122
chromium	249	224	2800 - 4270
cobalt	113	101	172 - 275
copper	94.9	85.5	82.0 - 120
Iron .	24400*	12500	70.4 - 100
lead	184	162	5480 - 19500
magnesium	3780*	2160	132 - 192
manganese	703		1650 - 2670
mercury	5.32	415	330 - 500
molybdenum		5.18	3.42 - 6.87
nickel	80.2 137	68.8	52.7 - 84.9
potassium		119	98.5 - 140
selenium	33000*	2840	2160 - 3520
silver	146	135	104 - 166
sodium	127	107	49.8 - 164
strontium	15600*	1010	709 - 1310
thallium	326	111	89.0 - 133
tin	106	99.3	76.8 - 122
	175	148	70.6 - 225
titanium	3100*	283	104 - 463
vanadium	151	104	70.5 - 138
zinc	311	275	222 - 328

# ADDENDUM SOP NO CHANGE FORM

### KATAHDIN ANALYTICAL SERVICES, INC. SOP "REVIEW WITH NO CHANGES" FORM

Name of Person Reviewing SOP: Nicholas Tillotson

Review Date: 2/2/12	
SOP Number: CAGOS	
SOP Title: 3050 DIG	
THE ABOVE REFERENCED SOP HAS BEEN REVIEWED ANALYST OR SUPERVISOR. NO CHANGES ARE REQU	
Department Supervisor Signature:	Date:
- Chamber	02/02/12
QAO Signature:	Date:
Leseic Dimond	020912

## KATAHDIN ANALYTICAL SERVICES, INC. SOP "REVIEW WITH NO CHANGES" FORM

Name of Person Reviewing SOP: Nicholas Tillotson

Review Date: 1/22/13	
SOP Number: CA - 605	
sop Title: Acid digestion of solid so	imples by USEPA
sop Title: Acid digestion of solid so method 3050 for metals analy	sis by ICP
THE ABOVE REFERENCED SOP HAS BEEN REVIEWED BY A QUALIFIED AND TRAINED ANALYST OR SUPERVISOR. NO CHANGES ARE REQUIRED TO THE SOP AT THIS TIME	
Department Supervisor Signature:	Date:
Meur	02/26/13
QAO Signature:	Date:
Liseie Dimond	02/26/13

## KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

Revision History:

SOP Number: CA-604 Revision History Cover Page Page 1

	George	Brewer		_Date:_	11/97
		4			
sor: _	Lloy	ge Bruce	1	Date:	01/19/01
nager: _	Joh C	Buter		Date:_	1/22/01
_	Dete	nah J. Ke	adeau	Date:_	1.22.01
ger: _	Der	nace f. W	efar	Date:_	1/22/01
		sor: Joh C	sor: Joh C. Butan  Quetorah J. M.	Sor: Joh C. Butan  Quetorah J. Madeau	nager:

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
01 3010A	Fermat changes, added pollution prevention, block digester; revised detabase references; vevised and added tables.	<i>On</i>	1:22:01	1/22/01
02	Added wording allowing use of digestates for ICP-MS and USis. Added use of block digester as primary heating source of adjusted volumes. Revised standard solution names of concs. in Figures 3044.	DN	8.29.02	8-29-03
03	Added Uranium to spiking socutions for LCS is MS/D. Removed the Internal Custody Record for Metals Digestates figure and reference.	LAN	04/06	04/06
04	Minor changes to Section 7 to reflect current practices. Updated Figure 1 - Sample Prep Logbook. Updated Figure 2 and 3 - Spike amounts.	LAN	05/09	05/09
05	Added references. Updated Figure 2 and 3 with correct spike information. Added CA-108 reference for subsampling information.	LAN	04/10	04/10

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TITLE:	ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS
	cknowledge receipt of this standard operating procedure by signing and dating both of the rovided. Return the bottom half of this sheet to the QA Department.
AQUEO	rledge receipt of copy of document SOP CA-604-05, titled ACID DIGESTION OF JS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL COLVED METALS.
Recipien	t:Date:
	IN ANALYTICAL SERVICES, INC. RD OPERATING PROCEDURE
AQUEO	rledge receipt of copy of document SOP CA-604-05, titled ACID DIGESTION OF JS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL COLVED METALS.
Recipien	t:Date:

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## TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

### 1.0 SCOPE AND APPLICATION

The purpose of this SOP is to describe the procedure utilized by Katahdin Analytical Services, Inc. personnel to solubilize metals in aqueous samples, wastes that contain suspended solids, and mobility-procedure extracts prior to analysis by inductively coupled plasma atomic emission spectroscopy (ICP) and inductively coupled plasma mass spectrometry (ICP-MS). This SOP applies to samples prepared by EPA Method 3010, with the method modifications mentioned in Table 2.

### 1.1 Definitions - none.

### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of analysts experienced in the acid digestion of aqueous samples by EPA Method 3010. Each analyst must demonstrate and document their ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, current revision, "Personnel Training & Documentation of Capability".

It is the responsibility of all Katahdin technical personnel involved in the acid digestion of aqueous samples using EPA Method 3010 to read and understand this SOP, to adhere to the procedures outlined, and to properly document their work in the appropriate lab notebook. Any deviations from the method or irregularities with the samples should also be recorded in the lab notebook and reported to the Supervisor or designated qualified data reviewer responsible for these data.

It is the responsibility of the Supervisor to ensure that technical personnel perform acid digestions in accordance with this SOP and to confirm that their work is properly documented through periodic review of the associated logbooks.

### 1.3 Safety

The acids used in this procedure are highly corrosive and reactive, and spiking standards contain toxic metals. The toxicity and reactivity of client samples are usually unknown, so samples should always be assumed to present a contact hazard. To reduce or eliminate exposure to potentially harmful chemicals, lab coats, gloves, and safety glasses or goggles must be worn whenever handling samples or reagents. Additional safety apparel, including face shields, rubber aprons, dust masks, and rubber shoe protectors, is available in the metals prep lab and should be worn whenever circumstances warrant.

Acids should be added to samples slowly and carefully while watching for reactions. This should be done under a hood, in case harmful fumes are evolved.

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## TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

Hood sashes should be lowered as far as possible whenever beakers are being heated in the hood. Use caution when handling hot beakers.

Each qualified analyst or technician must be familiar with Katahdin Analytical Health and Safety Manual including the Katahdin Hazardous Waste Management Plan and must follow appropriate procedures. These include the use of appropriate personal protective equipment (PPE) such as safety glasses, gloves and lab coats when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location of all safety equipment. Each analyst shall receive a safety orientation from their supervisor, or designee, appropriate for the job functions they will perform.

### 1.4 Pollution Prevention/Waste Disposal

Whenever possible, laboratory personnel should use pollution prevention techniques to address their waste generation. Refer to the current revision of the Katahdin Hazardous Waste Management Program for further details on pollution prevention techniques.

Excess spiking solutions must be emptied into the corrosive waste carboy located in the metals prep lab for subsequent appropriate disposal in accordance with the Chemical Hygiene Plan and Safety Manual.

Sample digestates should be stored for a minimum of 60 days after digestion to allow for analysis, and reanalysis if necessary. Digestates older than 60 days may be emptied into the corrosive waste carboy in the metals prep lab for subsequent appropriate disposal in accordance with the Chemical Hygiene Plan and Safety Manual.

Any other wastes generated during the preparation of samples must be disposed of in accordance with the Katahdin Chemical Hygiene Plan and Safety Manual and SOP SD-903, "Sample Disposal," current revision.

### 2.0 SUMMARY OF METHOD

The aqueous sample is refluxed with nitric acid in a covered digestion vessel. Additional nitric acid is added until the color of the digestate has stabilized. After the digestate has been evaporated to a low volume, it is refluxed with hydrochloric acid and diluted to the appropriate final volume with reagent water.

Samples may be concentrated (i.e. final digestate volume less than initial sample volume) during digestion if lower detection limits are required. Volumes of reagents and spiking standards must be added in proportion to the final volume of the digestate. Because concentration of samples during digestion increases the concentrations of dissolved solids

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## TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

and may exacerbate analytical interferences, concentration factors greater than 5 are not recommended.

### 3.0 INTERFERENCES

Interferences are discussed in the applicable analytical SOPs.

### 4.0 APPARATUS AND MATERIALS

- 4.1. 250 mL and 400 mL pre-cleaned Griffin beakers (cleaned according to the current revision of SOP CA-100, "Labware Cleaning") for digestion using a hot plate. If digestion will be performed using a block digester, 70ml graduated, polyethylene block digester tubes (with attached snap caps) will be used instead of glass beakers.
- 4.2 Ribbed watch glasses. If digestion is performed using a hot plate, 75 mm diameter and 100 mm diameter glass watch glasses (pre-cleaned as above) are used. If digestion is performed using a block digester, 40mm diameter disposable polyethylene watch glasses are used.
- 4.3 Adjustable volume automatic pipets covering the range from 10 uL to 1000 uL and disposable pipet tips; calibrated Finn pipets or Eppendorf pipets are acceptable.
- 4.4 Disposable graduated polystyrene specimen containers with pouring lips, 200 mL capacity.
- 4.5 Hot plate, block digester, or other heating source adjustable and capable of maintaining a temperature of 90-95 C. Hot plates must be numbered for easy identification.
- 4.6 Device for measuring hot plate temperature. This may consist of a heat-resistant 100ml beaker containing reagent water in which a thermometer is immersed. When using a block digester, a digestion tube containing reagent water in which a thermometer is immersed may be used. The temperature of one hot plate is measured each day, on a rotating basis. The hot plate identification number and the measured temperature are recorded on the sample preparation logbook sheet.
- 4.7 Plastic funnels, pre-cleaned as in Section 4.1.
- 4.8 Filter funnel holders, capable of suspending plastic funnels above disposable specimen containers.
- 4.9 Polyethylene wash bottles for dispensing reagent water and 5% HNO3.

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## TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

- 4.10 Filter paper, Whatman No. 41 or equivalent. Filters are acid-washed immediately prior to use as follows. Place a pre-cleaned funnel in the funnel holder and put a disposable plastic specimen container under the funnel to collect the rinsates. Place a folded filter in the funnel and rinse three times with approximate 10 mL volumes of 5% HNO3, making sure the entire surface of the filter is wetted each time and allowing each rinse to drain completely before continuing. Then rinse three times with approximate 25 mL volumes of reagent water. Discard the rinsates into the appropriate waste container. The acid-washed filter is now ready for use.
- 4.11 Polyethylene sample containers with screw caps or graduated polyethylene sample containers with attached snap lids, 125 mL capacity. These are not necessary when using the block digester since the final digestates are stored in the digestion tubes.
- 4.12 Repipetters (adjustable repeating pipetters with reservoirs) for dispensing concentrated nitric acid and 1:1 HCl.

### 5.0 REAGENTS

- 5.1 Concentrated nitric acid, HNO<sub>3</sub> trace metals grade.
- 5.2 Concentrated hydrochloric acid, HCl trace metals grade.
- 5.3 Reagent water water that meets the performance specifications of ASTM Type II water (ASTM D1193).
- 5.4 Hydrochloric acid, 1:1. Add a volume of concentrated hydrochloric acid to an equivalent volume of reagent water and swirl gently to mix.
- 5.5 Nitric acid, 5% v/v. Add 25 mL concentrated HNO<sub>3</sub> to 475 mL reagent water in a 500 mL wash bottle. Cap, point the dispensing tip into a sink, and shake gently to mix.
- 5.6 Multi-element spiking solutions (as listed in Figure 3).

### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

Samples to be analyzed for dissolved metals should be filtered through a 0.45 um membrane filter and preserved as soon as possible after collection. Samples to be analyzed for total metals should be preserved, unfiltered, as soon as possible after collection. Aqueous samples are preserved by acidification with nitric acid to a pH of <2.

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## TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

Please refer to Katahdin Analytical Services SOP CA-108, "Basic Laboratory Technique", current revision, for information on subsampling.

### 7.0 PROCEDURES

- 7.1 Prior to performing the digestion, make a list of the samples that are to be digested. Enter digestion information (Katahdin Sample Numbers, QC Batch ID, preparation date, analyst initials, etc.) into the ACCESS computer spreadsheet. Print out a copy of the spreadsheet. With a permamament marker, make sample labels and attach to the polyethylene sample containers that will contain the digestates.
- 7.2 If using glass beakers as the digestion vessels, submerge previously cleaned beakers three times into a 10% nitric acid bath, then rinse three times with reagent water. The polyethylene digestion tubes used in conjunction with the block digester do not require acid rinsing or precleaning. Label the digestion vessels with sample numbers.
- 7.3 If digestion is performed using a block digester, the sample aliquot may be measured in the digestion vessel using the graduations on the digestion tubes. Measure 50 ml of well-mixed sample into a 70 ml block digestion tube. A larger sample aliquot may be used (up to 250 mL) if concentration of the sample during digestion is desired. Sample volumes larger than 50 mL may be digested in 250 mL beakers. Measure aliquot of well-mixed sample into a graduated specimen cup and transfer into a properly cleaned 250 mL beaker. Sample volumes of more than 50ml may not be digested using the 70ml block digester tubes. The volumes of reagents and spiking solutions used must be adjusted in proportion to the final digestate volume. The reagent and spiking solution volumes listed below are based on a final volume of 50 mL.
- 7.4 Add spike solutions to matrix spike samples and laboratory control samples (refer to Figure 3 for spiking instructions).
- 7.5 Use a repipetter to add 1.5 mL of concentrated HNO3 (per 50 mL final volume) to the sample. Cover with a ribbed watch glass and place on heatsource. Heat cautiously, without boiling the sample, and evaporate to a low volume (10 15 mL).
  - <u>NOTE</u>: Do not allow any portion of the bottom of the digestion vessel to go dry during any part of the digestion. If a sample is allowed to go to dryness, low recoveries may result. Should this occur, discard the digestate and re-prepare the sample.
- 7.6 Cool the sample and add another 1.5 mL aliquot (per 50 mL final volume) of concentrated HNO3. Cover and resume heating, increasing the temperature until a gentle reflux action occurs.

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## TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

- 7.7 Continue heating, adding additional acid as necessary, until the digestate is light in color or does not change in appearance with continued refluxing.
- 7.8 Evaporate digestate to a low volume (10 15 mL).
- 7.9 Cool the sample and use a repipetter to add 5 mL (per 50 mL final volume) of 1:1 HCl. Cover the sample and resume heating, refluxing for an additional 15 minutes to dissolve any precipitate or residue resulting from evaporation.
- 7.10 Allow the sample to cool.
- 7.11 If the digestate contains visible particulate material, it must be filtered. Use a precleaned funnel and acid-rinsed filter paper to filter the digestate into a clean graduated plastic specimen container or block digester digestion tube. Using a wash bottle, rinse the digestion vessel with reagent water and add the rinsates to the filter apparatus. After all of the liquid in the filter has drained into the specimen container or digestion tube, thoroughly rinse the filter three times with small (5-10 mL) volumes of reagent water, allowing the liquid to drain completely after each rinse.

If the digestion was performed using hot plates and the digestate does not contain particulate material, simply decant the digestate into a clean graduated specimen container (or graduated sample container with attached snap lid), rinse the beaker with reagent water, and add the rinsates to the container.

If the digestion was performed using a block digester and the digestate contains no visible particulate material, the digestate may be brought to final volume and stored in the digestion tube without decanting or rinsing.

- 7.12 Using the graduations on the specimen container, snap-lid container or digestion tube, dilute to the required final volume with reagent water. If a specimen container has been used, transfer the contents to the corresponding labeled polyethylene sample bottle, cap the bottle, and discard the empty specimen container. If a snap-lid container or digestion tube has been used, close and secure the snap-lid. Shake the container gently to mix. The digestate is now ready for analysis.
- 7.13 Review the ACCESS computer spreadsheet for accuracy. If any information is incorrect, make the necessary changes to the computer spreadsheet and print out a corrected copy. Do not discard the original copy of the spreadsheet. Record (hand write) the sample bottle ID, reagent lot numbers, spiking information, initial and final volumes, hot plate ID and hot plate temperature in the appropriate spaces on the spreadsheet. Record any method deviations, irregularities with the samples, or other pertinent observations at the bottom of the page, and sign and date the spreadsheet. Bind all copies of the spreadsheet in the sample preparation log. An example sample preparation logbook page (ACCESS spreadsheet) is included as Figure 1.

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## TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

7.14 Place each batch of digestates in a box labeled with the QC Batch ID, and put the box of digestates in the metals digestates storage area.

7.15 A condensation of the procedure described above is included in this SOP as Table3. A controlled copy of this table may be posted in the metals preparation laboratory for reference by the analyst.

### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

- 8.1 At least one preparation blank for waters (PBW) is processed concurrently with each digestion batch of 20 or fewer samples, and is used to assess contamination resulting from the digestion procedure. The PBW consists of an aliquot of reagent water that is digested using the same reagents as those used to digest associated samples. The initial and final volumes of the PBW must be identical to those of the associated samples (i.e., if the associated samples were concentrated during digestion, the PBW must also be concentrated). Refer to the appropriate analytical SOP for PBW acceptance criteria and corrective actions.
- 8.2 At least one laboratory control sample for waters (LCSW) is processed concurrently with each digestion batch of 20 or fewer samples. The LCSW consists of an aliquot of reagent water that is spiked to contain all analytes of interest at known concentrations, and is digested using the same reagents as those used to digest associated samples. The initial and final volumes of the LCSW must be identical to those of the associated samples (i.e., if the associated samples were concentrated during digestion, the LCSW must also be concentrated). Directions for spiking the LCSW are contained in Figures 3 and 4. The measured analyte recoveries for the LCSW are used to assess digestion method performance. Refer to the appropriate analytical SOP for LCSW recovery acceptance criteria and corrective actions.
- 8.3 Matrix spiked samples are processed concurrently with each digestion batch at a minimum frequency of one per digestion batch. A matrix spike sample consists of an aliquot of a sample that is spiked with known amounts of all analytes of interest. Matrix spike recoveries are used to assess the effects of sample matrix on digestion and analysis performance. Directions for spiking matrix spike samples are contained in Figures 3 and 4. Refer to the appropriate analytical SOP for matrix spike recovery acceptance criteria and corrective actions.
- 8.4 Matrix spiked duplicate samples are processed concurrently with each digestion batch at a minimum frequency of one per digestion batch. Matrix spiked duplicate samples are used to assess the precision of the digestion and analysis methods. Refer to the appropriate analytical SOP for matrix spike duplicate precision acceptance criteria and corrective actions.

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## TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

<u>NOTE</u>: Clients may choose specific samples for matrix spike and matrix spike duplicate analysis; otherwise, the choice is left to the person performing the digestion. The sample volumes available may restrict the choice of samples used for matrix spike and duplicate digestion. Field blank samples should not be chosen for matrix spike and matrix spike duplicate analysis.

8.5 The quality control measures and frequencies described above are minimum requirements. They are summarized for reference in Table 1. Individual clients and analytical programs may impose additional QC requirements.

### 9.0 METHOD PERFORMANCE

Refer to the applicable analytical SOPs for method performance information.

### 10.0 APPLICABLE DOCUMENTS/REFERENCES

Test Methods for Evaluating Solid Waste, Physical/Chemical Methods, USEPA SW846, 3<sup>rd</sup> Edition, Final Updates I, II, IIA, IIB, III, IIIA, IIIB and IV, February 2007, Method 3010A.

Department of Defense Quality Systems Manual for Environmental Laboratories (DOD QSM), Version 4.1, 04/22/09.

The National Environmental Laboratory Accreditation Conference (NELAC) Standards, June 2003.

Katahdin SOP CA-101, Equipment Maintenance and Troubleshooting, current revision.

Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, current revision.

### LIST OF TABLES AND FIGURES

Table 1	QC Requirements
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- Table 2 Summary of Method Modifications
- Table 3 Procedure Condensation
- Figure 1 Example Page From Metals Sample Preparation Logbook
- Figure 2 Preparation of Matrix Spikes, LCSs, and Spiking Solutions: Method 3010
- Figure 3 Element Concentrations in Matrix Spikes, LCSs, and Spiking Solutions: Method 3010

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TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

# TABLE 1 QC REQUIREMENTS

Analytical Method	QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action
3010	Preparation Blank for Waters (PBW)	One per prep batch of 20 or fewer samples	Refer to analytical method	Refer to analytical method
	Laboratory Control Sample for Waters (LCSW)	One per prep batch of 20 or fewer samples	Refer to analytical method	Refer to analytical method
	Matrix Spike Sample	One per prep batch	Refer to analytical method	Refer to analytical method
	Matrix Spike Duplicate Sample	One per prep batch	Refer to analytical method	Refer to analytical method
	Demonstration of analyst proficiency; accuracy and precision	One time demonstration by each analyst performing the method	Must pass all applicable QC for method	Repeat analysis until able to perform passing QC; document successful performance in personal training file

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## TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

# TABLE 2 SUMMARY OF METHOD MODIFICATIONS

TOPIC	KATAHDIN SOP CA-604-05	EPA METHOD 3010, current revision
Apparatus/Materials	Disposable plastic specimen cup used to measure sample volume.	Graduated cylinder used to measure sample volume.
	2) Digestion performed in 250 mL, 400 mL Griffin beaker, or 70ml digestion tube to facilitate evaporation.	2) Digestion performed in 150 mL Griffin beaker.
	3) Ribbed watch glass used throughout digestion to reduce contamination.	3) Ribbed and non-ribbed watch glasses alternated in digestion.
Procedures	Digestate may be analyzed for antimony and silver.	Digestate may not be analyzed for antimony and silver.
	2) Sample aliquots larger or smaller than 100 mL may be used.	2) Requires sample aliquot of 100 mL.
	3) Sample evaporated to 10 - 15 mL.	3) Sample evaporated to 5 mL.

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TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

### TABLE 3

### PROCEDURE CONDENSATION: EPA METHOD 3010

- 1. If performing digestion on a hot plate, rinse glass beakers and ribbed watch glasses 3 times in acid bath. Then rinse beakers and watch glasses 3 times with reagent water. If performing digestion with block digester, polyethylene digestion tubes do not require precleaning.
- 2. Label digestion vessels with sample numbers.
- 3. Mix sample well, measure 50 mL (or smaller or larger aliquot) into a polyethylene digestion tube. If using glass beakers, measure aliquot into graduated specimen container, and transfer to appropriate digestion vessel.
- 4. Add spike solutions to matrix spike samples and LCSW (refer to Figure 3 of this SOP).
- 5. Add 1.5 mL (per 50 mL final volume) concentrated HNO3 to sample.
- 6. Cover with a ribbed watch glass.
- 7. Place on heating device (hotplate or block digester) and evaporate to 10 15 mL.
- 8. Cool sample and add another 1.5 mL (per 50 mL final volume) concentrated HNO3.
- 9. Resume heating until gentle reflux action occurs.
- 10. Continue heating, adding additional HNO<sub>3</sub> as necessary until digestion is complete.
- 11. Evaporate to 10 15 mL.
- 12. Cool sample and add 5 mL (per 50 mL final volume) 1:1 HCl. Resume heating and reflux gently for 15 minutes.
- 13. Cool sample and filter (if necessary) or decant into a graduated polyetheyne digestion tube. Rinse beaker with reagent water and filter or decant rinsate into specimen container.
- 14. Dilute to appropriate final volume with reagent water.
- 15. Cap sample container and shake gently to mix.

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## TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

### EXAMPLE PAGE FROM METALS SAMPLE PREPARATION LOGBOOK

JT Bake    S   Spike   LC	sgent Information: r HNO3: h 11002 4 SSSpiking Informati LPP-SPK-1 (ID/Vol) SPK-INT1 (ID/Vol) spK-INT2 (ID/Vol) spK-4 (ID/Vol):	ion: : MV 1202 : MV 1202 : MV 1202	4 /	0,05 0,5 0,05	_mL _mL _mL	Hot I Start	Time/To	ock ID 'emp.:_ emp. :_	: A 930 19: [Sec 195 195.: ALC 8	5_°C °C	Fisher Fil	ter Paper: <u> </u>	<b>Method</b>		
		Initial	Initial	Final	Final					Initial	Initial	Final	Final	1 m-woodman	73411727
ample ID	Batch ID	Wt/Vol	Units	Vol	Units	MX	Meth	Anal.	Date	Color	Clarity	Color	Clarity	Artifacts	Bottle
CSWAB011CW0	AB01ICW0	0.05	L	0.05	L	AQ	IC	AJB	02/01/2010	N/A	N/A	N/A	N/A		-
BWAB01ICW0	AB011CW0	1	L	-	L	AQ	IC	АЈВ	02/01/2010	N/A	N/A	N/A	N/A		<del>-</del> <del>-</del> <del>-</del> <del>-</del> +
D0405-001	AB01ICW0		L	-	L	AQ	IC	AJB	02/01/2010						
D0405-001P	AB011CW0		L	_	L	AQ	IC	AJB	02/01/2010	-	-	-			
3D0405-001S	AB011CW0		L	-	L	AQ	IC	AJB	02/01/2010					-	- +-
SD0405-002	AB011CW0		L	-	L	AQ	IC	AJB	02/01/2010						- +
SD0405-003	AB011CW0		L	_	L	AQ	IC	AJB	02/01/2010						- +-
SD0405-004	AB01ICW0		L	_	L	AQ	IC	AJB	02/01/2010		_				
SD0405-005	AB01ICW0		L		L	AQ	IC	AJB	02/01/2010						
SD0422-001	AB01ICW0		L	_	L	AQ	IC	AJB	02/01/2010						5-
SD0423-001	AB01ICW0		L		L	AQ	IC	AJB	02/01/2010						
SD0429-001	AB011CW0		L		L	AQ	IC	AJB	02/01/2010						
SD0429-002	AB01TCW0		L		L	AQ	IC	АЈВ	02/01/2010					-	
SD0455-001	AB011CW0		L		L	AQ	IC	AJB	02/01/2010		- 2			-	- +
SD0455-002	AB01ICW0		L		L	AQ	IC	АЈВ	02/01/2010						- +-
SD0455-003	AB01ICW0		L		L	AQ	IC	AJB	02/01/2010						- +-
SD0455-004	AB011CW0	3	L	1	L	AQ	IC	AJB	02/01/2010						
	2000										10.40				
								116.5							
				DOD		2-1	10_								
				on	_	JA.									
				DO											

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TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

### FIGURE 2

## PREPARATION OF MATRIX SPIKES, LABORATORY CONTROL SAMPLES, AND SPIKING SOLUTIONS FOR DIGESTION OF AQUEOUS SAMPLES BY METHOD 3010

Sample or Solution Name	Component Solution Name	Source of Component	Amount of Component Added per 50 mL Final Volume (mL)
	CLPP-SPK-1	Inorganic Ventures	0.050
Laboratory Control	CLPP-SPK-INT1	Lab Prepared (see below)	0.50
Sample (LCSW) and Matrix Spike	CLPP-SPK-INT2	Lab Prepared (see below)	0.50
	1000 mg/L Uranium Standard	Inorganic Ventures	0.005

Sample or Solution Name	Component Solution Name	Source of Component	Amount of Component Added per 100 mL Final Volume (mL)
	1000 mg/L Se	High Purity Standards	1.0
	1000 mg/L As	High Purity Standards	1.0
	1000 mg/L Pb	High Purity Standards	1.0
CLPP-SPK-INT1	1000 mg/L Cd	High Purity Standards	2.5
	1000 mg/L Sb	High Purity Standards	1.0
	10000 mg/L K	High Purity Standards	10.0
	10000 mg/L Na	High Purity Standards	7.5
	10000 mg/L Mg	High Purity Standards	5.0
	10000 mg/L Ca	High Purity Standards	2.5
	2007ICS-1	Inorganic Ventures	10.0
CLPP-SPK-INT2	1000 mg/L Sr	High Purity Standards	5.0
CLI I -SFK-INIZ	1000 mg/L Sn	High Purity Standards	5.0
	10000 mg/L Si	High Purity Standards	5.0

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TITLE: ACID DIGESTION OF AQUEOUS SAMPLES BY EPA METHOD 3010 FOR ICP AND ICP-MS ANALYSIS OF TOTAL OR DISSOLVED METALS

### FIGURE 3

ELEMENT CONCENTRATIONS IN MATRIX SPIKES, LABORATORY CONTROL SAMPLES, AND THEIR COMPONENT SPIKING SOLUTIONS FOR DIGESTION OF AQUEOUS SAMPLES BY METHOD 3010

		CONCENTRATION IN SOLUTION, mg/L										
Element	Matrix Spike	LCSW	CLPP- SPK-1	CLPP- SPK-4	CLPP- SPK- INT1	CLPP- SPK- INT2	2007 ICS-1	1000 mg/L U				
Aluminum	2.000	2.000	2000									
Antimony	0.500	0.500		100	100							
Arsenic	0.500	0.500		4	10							
Barium	2.000	2.000	2000									
Beryllium	0.050	0.050	50									
Boron	0.500	0.500		50		50	500					
Cadmium	0.250	0.250		5	25							
Calcium	2.500	2.500			250							
Chromium	0.200	0.200	200									
Cobalt	0.500	0.500	500									
Copper	0.250	0.250	250									
Iron	1.000	1.000	1000									
Lead	0.500	0.500		2	10							
Magnesium	5.000	5.000			500							
Manganese	0.500	0.500	500									
Molybdenum	0.300	0.300		30		30	300					
Nickel	0.500	0.500	500									
Potassium	10.000	10.000			1000							
Selenium	0.500	0.500		5	50							
Silicon	5.230	5.230				523	230					
Silver	0.050	0.050	50									
Sodium	7.500	7.500			750							
Strontium	0.500	0.500		50		50						
Thallium	0.500	0.500		5	10							
Tin	0.500	0.500		50		50						
Titanium	1.000	1.000		100		100	1000					
Uranium	0.100	0.100						1000				
Vanadium	0.500	0.500	500									
Zinc	0.500	0.500	500									

# ADDENDUM SOP NO CHANGE FORM

# KATAHDIN ANALYTICAL SERVICES, INC. SOP "REVIEW WITH NO CHANGES" FORM

Name of Person Reviewing SOP: Heather Henn	ingsen
Review Date: 5-1(-1)	
<b>SOP Number:</b> CA-602(-05	
SOP Title: Acid digestion of acqueaus samples ICP and ICP-MS analysis of total	by EPA method 3010 for .1 or dissolved metals.
THE ABOVE REFERENCED SOP HAS BEEN REVIEWED ANALYST OR SUPERVISOR. NO CHANGES ARE REQU	
Department Supervisor Signature:	Date:
- Story Mener	05/11/11
QAO Signature:	Date:
Leseis Dirnond	051111

# KATAHDIN ANALYTICAL SERVICES, INC. SOP "REVIEW WITH NO CHANGES" FORM

Name of Person Reviewing SOP: Nicholas Tillot	roz
Review Date: 1/18/12	
SOP Number: CA604-3010 D16	
SOP Title: SW3050B	
THE ABOVE REFERENCED SOP HAS BEEN REVIEWED ANALYST OR SUPERVISOR. NO CHANGES ARE REQU	day.
Department Supervisor Signature:	Date:
Y. Brewer	02/02/12
QAO Signature:	Date:
Jeseis Dimond	020912

# KATAHDIN ANALYTICAL SERVICES, INC. SOP "REVIEW WITH NO CHANGES" FORM

Name of Person Reviewing SOP: Nicholas Tillotson

Review Date: 1/22/13

SOP Number: CA-604 SOP Title: Acid digestion of EPA method 3010	aqueous samples by for ICP and ICP-MS
THE ABOVE REFERENCED SOP HAS BE ANALYST OR SUPERVISOR. NO CHANG	EEN REVIEWED BY A QUALIFIED AND TRAINED SES ARE REQUIRED TO THE SOP AT THIS TIME
Department Supervisor Signature:	Date:
- F. Drewer	02/26/13
QAO Signature:	Date:
_ Lislie Dimond	022613

## KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-627 Revision History Cover Page Page 1

TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020				
Prepared By:	George Binwer	Date:_	03/01	
Approved By:				
Group Supervisor:	- Glorge Breuzer	Date:_	04/02/01	
Operations Manage	r: Joh C. Benton	Date:_	3/29/01	
QA Officer:	Detorah J. Nadean	Date:_	03.27.01	
General Manager:	Derover Phulak	Date:_	04/03/01	
			* }	
Revision History:				

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
Ð١	Changed acid Solution Conc.  changed Run ID Naming convention added data reduction and reporting procedures  updated Standards tables (4-8)  updated Table 10 in include ISIS Configuration	LAD	07-16-05	02-16-05
<b>D</b> 2	sect. 4.2 - changed tubingsize  Sect. 5 - changed acid conc.s  Sect. 7 - major changes to reflect current practises including reporting data in the metals data-  losse. Sect. 8 - majorchanges updouting acceptance oritria. updated Tables 1,3-8,10 ; 11	LAY	04/06	04/06
03	Updated Tables 4.5 and 6 with corrent standards. Updated Table 1 with serial dilution, Post Digestion Metrix spike, MSA, ICS-A, ICS-AB and IDL mininum frequency or criteria. Updated Sect. 8 regarding Client specific requirements.	LAD	07/07	07/07
04	Section 7.18-changed instrument identifier to reflect new instrument; section 8-changed exceptance criteria and ICSAB analyte list; Table 1. whated acceptance criteria and corrective action to QC. Table 3-added all analytes to list-removed for information only "list.	LAD	04/08	04/08
05	updates to reflect changes from 6000 to 602A Added Handness by calculation attachment. Added LL OC requirement and criteria to Sect 8 and Table 1. Added criteria to analyze POL Gtd. at beginning and EUD of run.	(A)	02/09	02/09

SOP Number: CA-627 Revision History Cover Page (cont.)

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TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020

			<u> </u>	
SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
	Sect. 8 and QC Tables - Added DOD QSM			
96	references and criteria. Section10- Added references. Tables 4 > 7 - Added	LAD	08/09	08/09
	information pertaining to CCV conc. change	<u> </u>		
on ,	Adoled Table 2 with DoD QSM ver. 4.1 QC requirements. Updated Section 4.1, Table 10 und Table 11 with new autosampler information.	LAO	04/10	oulio
	Sect. 1.1 - Added autinitions. sect. 54.1, 42, 5.2			
08	7.9, 7.10, 7.1, 7.16 and 8.7. minor changes to reject current practice. Sect. 9 - added MDL, LODGARD LOD information. Sect 10- Madded, collited talterences. Updated Tables edited rejetences caro 042512	UAVO	04/12	04/12
	Added, tolled takenences. Updated Tables	1,5,6.7.	and 9	
	edited references caro 042512			
	·			

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TITLE:	TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020
	nowledge receipt of this standard operating procedure by signing and dating both of the vided. Return the bottom half of this sheet to the QA Department.
	ge receipt of copy of document SOP CA-627-08, titled TRACE METALS BY ICP-MS USING USEPA METHOD 6020
Recipient:	Date:
	ANALYTICAL SERVICES, INC. O OPERATING PROCEDURE
	ge receipt of copy of document SOP CA-627-08, titled TRACE METALS BY ICP-MS USING USEPA METHOD 6020
Recipient:	Date:

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TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020

### 1.0 SCOPE AND APPLICATION

Inductively coupled plasma-mass spectrometry (ICP-MS) is applicable to the determination of sub-ppb (ug/L) concentrations of a large number of elements in water samples and in waste extracts or digests. When dissolved constituents are required, samples must be filtered and acid-preserved prior to analysis. No digestion is required prior to analysis for dissolved elements in water samples. Acid digestion prior to filtration and analysis is required for groundwater, aqueous samples, industrial wastes, soils, sludges, sediments, and other solid wastes for which total (acid-leachable) elements are required.

ICP-MS has been applied to the determination of over 60 elements in various matrices. Analytes for which EPA has demonstrated the acceptability Method 6020 in a multi-laboratory study on solid wastes are listed as "analytes" in Table 4. Instrument detection limits, sensitivities, and linear ranges will vary with the matrices, and operating conditions. If Method 6020 is used to determine any analyte not listed in Table 4, it is the responsibility of the analyst to demonstrate the accuracy and precision of the method in the waste to be analyzed. The analyst is always required to monitor potential sources of interferences and take appropriate action to ensure data of known quality.

An appropriate internal standard is required for each analyte determined by ICP-MS. Recommended internal standards are <sup>6</sup>Li, <sup>45</sup>Sc, <sup>89</sup>Y, <sup>103</sup>Rh, <sup>115</sup>In, <sup>159</sup>Tb, <sup>165</sup>Ho, and <sup>209</sup>Bi. The lithium internal standard should have an enriched abundance of <sup>6</sup>Li, so that interference from lithium native to the sample is minimized. Other elements may need to be used as internal standards when samples contain significant amounts of the recommended internal standards.

### 1.1 Definitions:

<u>CCB</u> - Continuing Calibration Blank - An analyte-free solution consisting of acidified reagent water used to verify calibration accuracy periodically during analysis.

<u>CCV</u> - Continuing Calibration Verification - A midrange standard used to verify calibration accuracy periodically during analysis.

<u>Duplicate</u> - A second aliquot of a sample that is prepared and analyzed in the same way as the original sample in order to determine the precision of the method.

<u>ICB</u> - Initial Calibration Blank - An analyte-free solution consisting of acidified reagent water used to verify calibration accuracy.

<u>ICP-MS</u> - Inductively Coupled Plasma Mass Spectrometry.

<u>ICS</u> - Interference Check Samples - Two standards (ICS-A and ICS-AB) used to verify the effectiveness of interference correction equations. Solution ICS-A contains only interferents (AI, Ca, Fe, Mg, Na, K, P, S, Mo, Ti, C, CI) at high

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### TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020

concentrations; solution ICS-AB contains interferents at the same concentrations as well as analytes at low (20 ug/L) concentrations.

<u>ICV</u> - Initial Calibration Verification - A standard made from a source independent from the calibration standards and with analyte concentrations different from those in the CCV; used to verify the accuracy of the instrument calibration.

<u>IDL</u> - Instrument Detection Limit - The lowest concentration of an analyte that can be determined with 95% confidence.

<u>Internal Standard</u> - Pure analytes added to a sample, extract, or standard solution in known amounts and used to measure the relative responses of other method analytes that are components of the same sample or solution. Internal standards must be analytes that are not native to the sample.

- <u>LCS</u> Laboratory Control Sample A standard or solid reference material that has been brought through the sample preparation process.
- <u>LDR</u> Linear Dynamic Range The concentration range over which the instrument response to an analyte is linear.
- <u>LOD</u> Limit of Detection An estimate of the minimum amount of a substance that an analytical process can reliably detect. An LOD is analyte and matrix-specific and is used for DoD QSM acceptance criteria.
- <u>LOQ</u> Limit of Quantitation.- The minimum concentration of a target analyte that produces a quantitative result within specified limits of precision and bias.
- <u>PB</u> Preparation Blank Reagent water that has been brought through the sample preparation process.

<u>Post-Digestion</u> <u>Spike</u> - An aliquot of a sample to which a known amount of analyte has been added before analysis and after digestion, if digestion is required.

<u>PQL</u> - Practical Quantitation Limit - The lowest concentration of an analyte that is routinely reported by the laboratory; nominally three to five times the IDL.

<u>Matrix Spike</u> - An aliquot of a sample to which a known amount of analyte has been added before digestion.

<u>Serial Dilution</u> - The dilution of a sample by a factor of five. When corrected by the dilution factor, the measured analyte concentrations of the diluted sample should agree with those of the original undiluted sample within specified limits. Serial dilution may reflect the influence of interferents.

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### TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020

### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of, analysts experienced in ICP-MS analysis by USEPA Method 6020 who are knowledgeable in the recognition and in the correction of spectral, chemical, and physical interferences in ICP-MS. Each analyst must demonstrate and document their ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, current revision, "Personnel Training & Documentation of Capability".

It is the responsibility of all Katahdin technical personnel involved in ICP-MS analysis by USEPA Method 6020 to read and understand this SOP, to adhere to the procedures outlined, and to properly document their data in the appropriate lab notebook. Any deviations from the test or irregularities with the samples should also be recorded in the lab notebook and reported to the Department Manager or designated qualified data reviewer responsible for this data.

It is the responsibility of the Department Manager to oversee that members of their group follow this SOP, to ensure that their work is properly documented, and to initiate periodic review of the associated logbooks.

### 1.3 Safety

Users of this procedure must be cognizant of inherent laboratory hazards, proper disposal procedures for contaminated materials and appropriate segregation of hazardous wastes. The toxicity or carcinogenicity of each reagent used in this method has not been precisely defined; however, each chemical should be treated as a potential health hazard. A reference file of material safety data sheets is available to all personnel involved in the chemical analysis. Everyone involved with the procedure must be familiar with the MSDSs for all the materials used in this procedure.

Each qualified analyst or technician must be familiar with Katahdin Analytical Environmental Health and Safety Manual including the Katahdin Hazardous Waste Plan and must follow appropriate procedures. These include the use of appropriate personal protective equipment (PPE) such as safety glasses, gloves and lab coats when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location of all safety equipment. Each analyst shall receive a safety orientation from their Department Manager, or designee, appropriate for the job functions they will perform.

Samples, sample digestates, standards, and other reagents used in ICP analysis may contain high concentrations of acids and toxic metals. Spilled samples and reagents should be cleaned up from instrument and laboratory surfaces immediately.

Liquid argon represents a potential cryogenic and suffocation hazard and safe handling procedures should be employed at all times when handling liquid argon

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### TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020

tanks and fittings. Safety glasses and cryogenic-resistant gloves should be worn when changing or adjusting argon tanks.

The Agilent 7500 ICP-MS spectrometer is safety-interlocked to prevent user exposure to harmful electrical voltages, radio frequency emissions, ultraviolet radiation, high temperatures, and other hazards. At no time should the operator attempt to disable these interlocks or operate the instrument if any safety interlock is suspected to be disabled

### 1.4 Pollution Prevention/Waste Disposal

Whenever possible, laboratory personnel should use pollution prevention and waste minimization techniques to address their waste generation. Refer to the current revision of the Katahdin Hazardous Waste Management Program for further details on pollution prevention techniques.

Samples, sample digestates, standards, and other reagents used in ICP-MS spectrometry may contain high concentrations of acids and toxic metals. They should be disposed of in a manner appropriate to the types of hazards they present. All digested samples and excess reagents and standards should be disposed of in the satellite waste container for corrosive wastes (labeled "Waste Stream A") that is located in the Metals Instrument lab. Further information regarding waste classification and disposal may be obtained by consulting Katahdin Analytical Environmental Health and Safety Manual and the Department Manager.

### 2.0 SUMMARY OF METHOD

- 2.1 Prior to analysis, samples that require total ("acid-leachable") values must be digested using appropriate sample preparation methods (such as USEPA Methods 3005 3051).
- USEPA Method 6020 describes the multi-elemental determination of analytes by ICP-MS. The method measures ions produced by a radio-frequency inductively coupled argon plasma. Analyte species originating in a liquid are nebulized and the resulting aerosol transported by argon gas into the plasma torch. The ions produced are entrained in the plasma gas and introduced, by means of a vacuum interface, into a mass spectrometer. The ions produced in the plasma are sorted according to their mass-to-charge ratios and quantified with a channel electron multiplier. Interferences must be assessed and valid corrections applied or the data flagged to indicate problems. Interference correction must include compensation for background ions contributed by the plasma gas, reagents, and constituents of the sample matrix.

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### 3.0 INTERFERENCES

Isobaric elemental interferences in ICP-MS are caused by isotopes of different elements forming atomic ions with the same nominal mass-to-charge ratio (m/z). The Agilent 7500 ChemStation data system is used to correct for these interferences. This involves determining the signal for another isotope of the interfering element and subtracting the appropriate signal from the analyte isotope signal. Isobaric molecular and doubly-charged ion interferences in ICP-MS are caused by ions consisting of more than one atom or charge, respectively. Most isobaric interferences which could affect ICP-MS determinations have been identified. Examples include ArCl<sup>+</sup> ions on the As signal and MoO<sup>+</sup> ions on the cadmium isotopes. While the approach used to correct for molecular isobaric interferences is demonstrated below using the natural isotopic abundances from the literature, the most precise coefficients for an instrument must be determined from the ratio of the net isotope signals observed for a standard solution at a concentration providing suitable (<1 percent) counting statistics. Because the <sup>35</sup>Cl natural abundance of 75.77 percent is 3.13 times the <sup>37</sup>Cl abundance of 24.23 percent, the chloride correction for arsenic can be calculated (approximately) as follows (where the <sup>38</sup>Ar<sup>37</sup>Cl<sup>+</sup> contribution at m/z 75 is a negligible 0.06 percent of the <sup>40</sup>Ar <sup>35</sup>Cl<sup>+</sup> signal):

Corrected <sup>75</sup>As signal (using natural isotopic abundances for coefficient approximations) =

(m/z 75 signal) - (3.13) (m/z 77 signal) + (2.73) (m/z 82 signal),

where the final term adjusts for any selenium contribution at 77 m/z.

<u>NOTE:</u> Arsenic values can be biased high by this type of equation when the net signal at m/z 82 is caused by ions other than  $^{82}$ Se $^{+}$ , (e.g.,  $^{81}$ BrH $^{+}$  from bromine wastes or  $^{82}$ Kr from krypton contamination in the Ar). Similarly:

Corrected <sup>114</sup>Cd signal (using natural isotopic abundances for coefficient approximations) = (m/z 114 signal) - (0.027) (m/z 118 signal) - (1.63)(m/z 108 signal),

where last 2 terms adjust for any tin or MoO<sup>+</sup> contributions at m/z 114.

<u>NOTE:</u> Cadmium values will be biased low by this type of equation when <sup>92</sup>ZrO<sup>+</sup> ions contribute at m/z 108. Also, use of m/z 111 for Cd is even subject to direct (<sup>92</sup>ZrOH<sup>+</sup>) ions and indirect (<sup>90</sup>ZrO<sup>+</sup>) additive interferences when Zr is present.

<u>NOTE:</u> As for the arsenic equation above, the coefficients in the Cd equation are only illustrative. The most appropriate coefficients for an instrument can be determined from the ratio of the net isotope signals observed for a standard solution at a concentration providing suitable (<| percent) counting precision.

The interference correction equations that are used by this laboratory in performing USEPA Method 6020 are listed in Table 4. The accuracy of these types of equations is based upon

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### TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020

the constancy of the observed isotopic ratios for the interfering species. Corrections that presume a constant fraction of a molecular ion relative to the "parent" ion have not been found to be reliable, e.g., oxide levels can vary. If a correction for an oxide ion is based upon the ratio of parent-to-oxide ion intensities, the correction must be adjusted for the degree of oxide formation by the use of an appropriate oxide internal standard previously demonstrated to form a similar level of oxide as the interferent. This type of correction has been reported for oxide-ion corrections using ThO+/Th+ for the determination of rare earth elements. The use of aerosol desolvation and/or mixed plasmas have been shown to greatly reduce molecular interferences (the Agilent 7500 ICP-MS spectrometer employs spray chamber cooling to effect aerosol desolvation). These techniques can be used provided that method detection limit, accuracy, and precision requirements for analysis of the samples can be met.

- 3.1 Physical interferences are associated with the sample nebulization and transport processes as well as with ion-transmission efficiencies. Nebulization and transport processes can be affected if a matrix component causes a change in surface tension or viscosity. Changes in matrix composition can cause significant signal suppression or enhancement. Dissolved solids can deposit on the nebulizer tip of a pneumatic nebulizer and on the interface skimmers (reducing the orifice size and the instrument performance). Total solid levels below 0.2% (2,000 mg/L) are recommended to minimize solid deposition. An internal standard can be used to correct for physical interferences, if it is carefully matched to the analyte so that the two elements are similarly affected by matrix changes. The internal standard used should differ from the analyte of interest by no more than 50 amu. See table 14 for a list of internal standards used. When the intensity level of an internal standard is less than 70 percent or greater than 120 percent of the intensity of the first standard used during calibration, the sample must be reanalyzed after a fivefold (1+4) or greater dilution has been performed.
- 3.2 Memory interferences can occur when there are large concentration differences between samples or standards that are analyzed sequentially. Sample deposition on the sampler and skimmer cones, spray chamber design, and the type of nebulizer affect the extent of the memory interferences that are observed. The rinse period between samples must be long enough to eliminate significant memory interference.

### 4.0 APPARATUS AND MATERIALS

4.1 Agilent 7500 ICP-MS system, consisting of the Agilent 7500 ICP-mass spectrometer and its controlling computer data station. The spectrometer is capable of providing resolution better than or equal to unit resolution at 10% peak height. The Agilent 7500 mass range of 2-260 amu exceeds the method requirement of 2- 240 amu. The Agilent 7500 ChemStation software allows automatic corrections for isobaric interferences and correction for internal standard responses as required by the method. All critical argon flows including nebulizer argon are under mass flow

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### TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020

controller control and a peristaltic pump is used for sample introduction. Peripheral equipment includes a Elemental Scientific SC-4 PX Fast Autosampler and Sample Introduction system, and Bullzip PDF printer set to print to file ICPMS\_CP.pdf located in folder PDF\_PRINTS on the desktop.

- 4.2 Peristaltic pump tubing 2-stop ESI PVC flared black-black (0.76 mm ID) and orange-green (0.38 mm ID). 3-stop Pharmed blue-yellow (1.52 mm ID).
- 4.3 15 ml 17x100 mm polypropylene or polystyrene disposable test tubes for samples and 50 ml polypropylene centrifuge tubes for standards.
- 4.4 Automatic adjustable-volume pipetters of suitable precision and accuracy. Calibrated Eppendorf Reference pipets and Finn digital pipets are appropriate.
- 4.5 Trace metal grade pipette tips.
- 4.6 Volumetric glassware or plasticware of suitable precision and accuracy.
- 4.7 Talc free vinyl gloves.
- 4.8 Argon gas supply (high purity grade gas or liquid, 99.99%).
- 4.9 For the determination of trace levels of elements, contamination and loss are of prime consideration. Potential contamination sources include improperly cleaned laboratory apparatus and general contamination within the laboratory environment from dust etc. A clean laboratory work area, designed for trace element sample handling must be used. Standards, samples and blanks should be exposed to the laboratory environment as little as possible. The use of preparation blanks and spikes should be used to verify the absence of sources of contamination and loss. If necessary, polypropylene sample tubes should be rinsed and stored in dilute acid prior to use.

<u>NOTE:</u> Chromic acid must not be used for cleaning glassware for trace metals analysis.

### 5.0 REAGENTS

Acids used in the preparation of standards and for sample processing must be of high purity. Redistilled acids are recommended because of the high sensitivity of ICP-MS. Mallincrodt/Baker "Instra-Analyzed" trace-metals grade acids are appropriate. It is important to match the acid concentration in standards and samples. Concentrations of antimony and silver between 50-500 ug/L require 1% (v/v) HCI for stability; for concentrations above 500 ug/L additional HCI will be needed. For this reason, it is recommended that antimony and silver concentrations in samples and standards be maintained below 500 ppb wherever possible. Acids

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### TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020

are received in poly-coated glass bottles, and are stored under the hood in the Metals sample preparation laboratory at room temperature until use. All acids are considered to have a shelf life of three years from date of receipt unless otherwise indicated by the vendor. Refer to the current revision of Katahdin SOP CA-105, Reagent and Solvent Handling, for further details.

- 5.2 Laboratory reagent grade water, trace metals free, equivalent to ASTM Type 1 (ASTM D 1193), >18 Megohm/centimeter resistivity.
- 5.3 Single element and multielement stock standard solutions purchased standards prepared from high purity salts or metals, and supplied by the vendors with certificates of purity and analysis. Refer to Tables 5 and 6 for a listing of stock standards required, and to Table 9 for element concentrations in stock standards. Purchased stock standards are received in polyethylene containers and are stored in their original containers at room temperature in the Metals Standards Preparation Laboratory. All purchased stock standards are given an expiration date as indicated by the manufacturer. Refer to the current revision of Katahdin SOP CA-106, Standard Preparation, Documentation and Traceability, for further details.
- Intermediate standard solutions laboratory-prepared multielement standards that are used in the subsequent preparation of working standards. Refer to Table 6 for a listing of intermediate standards required and for preparation instructions. Refer to Table 8 for element concentrations in intermediate standards. Intermediate standards are stored at room temperature in acid-washed polyethylene containers in the Metals Standards Preparation Laboratory. Intermediate standards are assigned an expiration date of three months from the date of preparation, or the earliest expiration date of a component standard, whichever comes first. Refer to the current revision of Katahdin SOP CA-106, Standard Preparation, Documentation and Traceability, for further details.
- 5.5 Working standard solutions - laboratory-prepared multielement standards that are used to calibrate the instrument, to provide internal standardization through on-line addition, and to perform all necessary QC checks. Refer to Table 5 for a listing of working standards and for preparation instructions. Refer to Table 7 for element concentrations in working standards. Working standards are stored at room temperature in acid-washed polyethylene containers in the Metals Standards Preparation Laboratory. All working standards except the ICSA and ICSAB solutions are assigned an expiration date of three months from the date of preparation, or the earliest expiration date of a component standard, whichever comes first. The ICSA and ICSAB solutions are assigned an expiration date of one week from the date of preparation, or the earliest expiration date of a component standard, whichever comes first. Refer to the current revision of Katahdin SOP CA-106, Standard Preparation, Documentation and Traceability, for further details.
- 5.6 Blanks: Three types of blanks are required for the analysis. The calibration blank is used in establishing the calibration curve. The preparation blank is used to monitor

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for possible contamination resulting from the sample preparation procedure. The rinse blank is used to flush the system between all samples and standards.

- 5.6.1 The calibration blank consists of the same concentrations of the same acid(s) used to prepare the final dilution of the analyte calibration solutions (currently 1% HNO<sub>3</sub> and 0.5% HCl, v/v, in laboratory reagent grade water). Use of HCl for antimony and silver is cited in Section 5.1.
- 5.6.2 The preparation blank must be carried through the complete preparation procedure and contain the same volumes of reagents as the associated digested sample solutions.
- 5.6.3 The rinse blank consists of 4% HNO<sub>3</sub> and 0.5% HCl,v/v, in reagent water. Prepare a sufficient quantity to flush the system between standards and samples.

### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

Samples to be analyzed for trace metals by ICP-MS should be collected and preserved as described in the following table.

Matrix	Container <sup>1</sup>	Collection Volume/ Weight	Preservation/ Treatment	Holding Time
Aqueous (total)	P, G	250 mL	HNO <sub>3</sub> to pH < 2	6 months
Aqueous	P, G	250 mL	Filter, HNO <sub>3</sub> to pH <	6 months
(dissolved)			2	
Solid	P, G	10 g	Cool, 4°C	6 months

<sup>&</sup>lt;sup>1</sup> P = polyethylene or, G = glass

### 7.0 PROCEDURES

- 7.1 Instrument control and data acquisition are completely automated through the use of the Agilent Chemstation software. The main Chemstation screen is accessed by double-clicking the "ICP-MS Top" icon on the Windows desktop. Autosampler tables are edited by selecting "Edit Sample Log Table" from the Sequence menu in the Agilent Chemstation software. In the following discussion, software menu items that are to be selected are printed in boldface. The instrument operating conditions, acquisition parameters, acquisition masses, and internal standards for analysis USEPA Method 6020 are detailed in Table 11.
- 7.2 Turn on the argon supply at the tank and set the pressure to >700 kPa.

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- 7.3 Turn on the water chiller/recirculator.
- 7.4 Verify that the exhaust hood is in operation.
- 7.5 Ensure that the internal standard solution bottle is adequately full. Consumption is approximately 2.5 mL/hour.
- 7.6 Verify that the rinse station reservoir has an adequate supply of reagent water.
- 7.7 Verify that the drain reservoir has adequate room to accept the day's drain waste. Empty the reservoir as necessary into an appropriate waste container (Waste Stream A) located in the Hazardous Waste Storage Area.
- 7.8 Inspect the peristaltic pump tubes for signs of flattening and wear, and replace them as necessary. Clamp the peristaltic pump tubes into the peristaltic pump.
- 7.9 Open the Chemstation software by double-clicking the "ICP-MS Top" icon. Initiate the plasma by selecting Instrument>>Instrument Control>>Plasma>>Plasma On and allow the instrument to aspirate calibration blank solution for at least 45 minutes. During this warm-up, select Tune>>Sensitivity>>Start to start the instrument scanning the mass range. Verify that the flow of sample and internal standard solutions through the uptake lines and into the nebulizer is free from pulsations by introducing an air bubble into each line and observing its progress. Adjust the pump clamp tension on each line to obtain a constant, pulse-free flow.
- 7.10 After the 45 minute warm-up, check the responses of masses 82 and 83 to insure minimal krypton intereference with selenium. Mass 83 response should be < 2000 counts per second. Then aspirate the Instrument Tune Solution (10 ppb Li, Y, Ce, Tl) and check the responses and RSDs at masses 7, 89, and 205.
- 7.11 Generate a tune report by selecting **Tune>>File>>Generate Report**. Evaluate the tune report against the tune specifications listed in Table 12. If the tune passes all specifications, proceed to step 7.14.
- 7.12 If the tune report indicates unacceptable instrument performance for any parameter, initiate an autotune by selecting **Tune>>Autotune>>Run**. The Chemstation software will attempt to tune the instrument to meet the tune specifications, and will generate a new tune report after autotuning. Evaluate the new tune report against the tune specifications listed in Table 12.
- 7.13 Repeat step 7.12 until all tune specifications have been met. File the final tune report.
- 7.14 Aspirate the P/A tuning solution (see Table6) and run a P/A autotune by selecting **Tune>>Tune>>P/A Factor>>Run**. This will calibrate the pulse and analog modes of the detector. File the P/A report with the Tune report.

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7.15 Load the instrument analytical method and calibrations table for USEPA Method 6020 into memory by selecting **Methods>>Load>>K6020.M>>K6020.C**.

- 7.16 Edit the sequence template "K6020.S" to create an analytical sequence table listing all of the samples to be analyzed. To do this, select "Edit Sample Log Table" from the **Sequence** menu in the Agilent Chemstation software. Double-click **SMPL** from the menu at the top left. Fill in the sample table with sample IDs, vial numbers, analytical method (K6020.M for all samples), dilution factors, and failure actions. When the sample table is complete, select **Print** to print this table. Close the "Edit Sample Log Table" window. Save the sample log table under a new name by selecting **Save** under the **Sequence** menu and then typing the name.
- 7.17 Load the autosampler racks according to the analytical sequence printout.
- 7.18 Select Sequence>>Load and Run Sequence, and select the appropriate autosampler sequence table from the displayed list. Enter the analyst's initials in the Operator box. Change data file name to appropriate designation. The protocol for naming data files is as follows: the 1st character is a letter that identifies the instrument ("J" for the Agilent 75000 ICP-MS), the 2<sup>nd</sup> character is a letter that identifies the year of the run ("X" for 2007, "Y" for 2008, etc.), the 3<sup>rd</sup> character is a letter that identifies the month of the run ("A" for January, "B" for February, etc.), the 4<sup>th</sup> and 5<sup>th</sup> characters are numbers that identify the date of the run ("01" for the first day of the month, etc.), and the 6<sup>th</sup> character is a letter that sequentially identifies the run ("A" for the first run of the day on that instrument, "B" for the second run, etc.). For example, the run identified as "JYA16A" is the first run of the day that was performed on January 16, 2008, using the Agilent 7500 ICP-MS. Select Run. The instrument will analyze all samples in the order listed in the table. Analysis must proceed in the sequence described in Table 10 to ensure that all necessary quality control samples are analyzed at the appropriate frequencies. A minimum of three replicate scans is required for all standards and samples. Analysis always begins with the analysis of a calibration blank followed by at least three multielement calibration standards to calibrate the instrument. The system is flushed with rinse blank between each sample and standard, and each sample and standard is aspirated for at least one minute prior to the beginning of mass scanning.
- 7.19 Analysis continues with analysis of the initial calibration verification standard (ICV) and the initial calibration blank (ICB) to verify the accuracy of the calibration. Refer to Section 8 and Table 1 for additional information.
- 7.20 A practical quantitation limit standard (PQL) is analyzed at the beginning of the run to verify calibration accuracy at the reporting limit. Refer to Section 8 and Table 1 for additional information.
- 7.21 A continuing calibration verification standard (CCV) and a continuing calibration blank (CCB) must be analyzed at the beginning of the run, after every ten samples,

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and at the end of the run to verify the continued accuracy of the calibration. Refer to Section 8 and Table 1 for additional information.

- 7.22 Interference check standard solutions ICS-A and ICS-AB must be analyzed at the beginning of each run and every 12 hours to verify the adequacy of interference corrections. Refer to Section 8 and Table 1 for additional information.
- 7.23 All sample analytical results for a particular element that are bracketed (preceded or followed) by failing results in a calibration verification sample (ICV, ICB, CCV, or CCB) for that element must not be reported, except as noted in Sections 8.5, 8.6, and 8.9. The sample must be reanalyzed for the element in question.
- 7.24 All samples that exceed the linear dynamic range must be diluted and reanalyzed.
- 7.25 If dilutions of digested samples are performed, the measured element concentrations must be multiplied by the dilution factor prior to reporting. This is accomplished automatically by entering the dilution factor in the sample log table prior to initiation of analysis.
- 7.26 If an element has more than one monitored isotope, examination of the concentration calculated for each isotope, or the isotope ratios, will provide useful information for the analyst in detecting a possible spectral interference. Consideration should therefore be given to both primary and secondary isotopes in the evaluation of the element concentration. In some cases, secondary isotopes may be less sensitive or more prone to interferences than the primary recommended isotopes, therefore differences between the results do not necessarily indicate a problem with data calculated for the primary isotopes. In the case of Pb, quantitation is based on the sum of isotopes 206, 207 and 208 to compensate for any variation in naturally occurring isotope ratios. This is accomplished through the use of the interference correction equation for lead.
- 7.27 Calculations, aqueous samples: Final element concentrations for aqueous samples are reported in units of micrograms per liter (ug/L). The reported concentrations are calculated from measured digestate concentrations as follows:

Concentration (ug/L) = 
$$\frac{MC \times DF \times FV}{IV}$$

where: MC = Measured digestate concentration (ug/L)

DF = Instrument dilution factor FV = Final digestate volume (L) IV = Digested sample volume (L)

7.28 Calculations, solid samples: Final element concentrations for solid samples are reported in units of milligrams per kilogram (mg/kg) on a dry weight basis. The

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reported concentrations are calculated from measured digestate concentrations as follows:

Concentration (mg/kg dry weight) =  $\frac{MC \times DF \times FV \times 100}{W \times S}$ 

where: MC = Measured digestate concentration (ug/L)

DF = Instrument dilution factor FV = Final digestate volume (L)

W = Weight of digested wet sample (g)

S = Percent solids

### DATA REDUCTION AND REPORTING

- 7.29 When the analytical run is completed, the analyst should print a run summary and create a data import file.
- 7.30 Follow these steps to print the run summary: Open the FileView program using the "FIVIEW" icon on the Windows Desktop. Select the data file of interest and move the required samples into the "Process List". Make sure the "Corrected Data" box is checked. Click the "Process" button. The data will be displayed in a spreadsheet format.
- 7.31 Select "Configure" from the top menu and "Sublists" from the displayed options. Select "Load Sublist" and then select "K6020" from this list of options. Make sure the "Enable Sublist" box is checked. Click the "OK" button. This will display only the analyte masses in the spreadsheet.
- 7.32 Select "Quant Info" from the top menu and "Quant Results" from the displayed options. This will display the data in concentration units. Click on the cell in the top left corner of the spreadsheet to highlight all data. Select "Tools" from the top menu and "Import Data into Spreadsheet Application" from the displayed options. Click the "Save" button. The data is transferred to a Microsoft Excel spreadsheet. Minimize this window and return to the FileView spreadsheet.
- 7.33 Select "Configure" from the top menu and "Sublists" from the displayed options. Select "Load Sublist" and then select "INTSTDS" from this list of options. Make sure the "Enable Sublist" box is checked. Click the "OK" button. This will display only the internal standard masses in the spreadsheet.
- 7.34 Select "Count Info" from the top menu and "Counts/sec" from this list of options. This will display the data in counts per second units. Click on the cell in the top left corner of the spreadsheet to highlight all data. Select "Tools" from the top menu and "Import Data into Spreadsheet Application" from the displayed options. Click the "Save" button. The data is transferred to a Microsoft Excel spreadsheet. Copy the internal standard names and results and paste into the analyte results Excel spreadsheet.

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- 7.35 In a separate cell of the Excel spreadsheet, input a formula to calculate the percent recovery of the internal standard relative to that of the calibration blank for each internal standard measurement of each sample.
- 7.36 Input a formula into the sample results cells to correct the data for dilution.
- 7.37 Format, print and save (as FileName.XLS, e.g. JYA28A.XLS) this run summary.
- 7.38 Follow these steps to create the data import file: Open the FileViewNT program using the "FIVIEWNT" icon on the Windows Desktop. Select the data file of interest and move the required samples into the "Process List". Make sure the "Corrected Data" box is checked. Click the "Process" button. The data will be displayed in a spreadsheet format.
- 7.39 Select "Configure" from the top menu and "Sublists" from the displayed options. Select "Load Sublist" and then select "K6020" from this list of options. Make sure the "Enable Sublist" box is checked. Click the "OK" button. This will display only the analyte masses in the spreadsheet.
- 7.40 Select "Quant Info" from the top menu and "Quant Results" from the displayed options. This will display the data in concentration units.
- 7.41 Select the "Transpose" from the menu.
- 7.42 Select "Tools" from the top menu and "Copy Selected Data to CSV File" from this list of options. Set the name the file as "FileName.CSV", e.g., "JYA28A.CSV". Click the "Save" button.

#### To import data into the Metals Database:

- 7.43 Select the "ICPMS Import" icon from the Windows Desktop, the ICPMS Import window will appear. Enter the datafile name without extension, (e.g., "JYA28A") and click the "Import" button.
- 7.44 When the "Import finished" message appears, close the ICPMS Import window and select the "KIMS\_METALS" icon from the Windows Desktop. The Metals Database Main Menu will appear.
- 7.45 Select the "Reporting Menu" button. From the Reporting Menu screen select the Batch QC Menu button and then the "Calculate Batch QC" button.
- 7.46 From the resulting list of QC results, deselect any items that fail run QC or do not need ICP-MS analysis. Click on the "Accept Selected Batch QC" button.
- 7.47 From the Metals Main Menu, select the "Additional Data Handling" button. The Data Menu will appear. Select the "Report Added Compounds" button.

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7.48 From the resulting list of sample results, deselect any items that fail run QC or do not need ICP-MS analysis. Click on the "Accept Data" button.

- 7.49 Once all associated data from an analysis run have been processed, go to the Generate Coverage portion of the Export Menu and print the Run Log and Logbook Page for the analysis run.
- 7.50 Combine these reports with the raw data printout and the Run Info Sheet and scan into a PDF format. Save in the "ICPMS DATA" section of the "METPDF" directory on the IMAGESERVER.

#### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

USEPA Method 6020 requires the laboratory to perform specific quality control checks to assess laboratory performance and data quality. Minimum frequencies, acceptance criteria, and corrective actions for these control checks are tabulated in Table 1 and are described below. Table 1 criteria are intended to be guidelines for analysts. The table does not cover all possible situations. If any of the QC requirements are outside the recovery ranges listed in Table 1, all associated samples must be evaluated against all the QC. In some cases data may be reported, but may be reanalyzed in other cases. Making new reagents and standards may be necessary if the standardization is suspect. The corrective actions listed in Table 1 may rely on analyst experience to make sound scientific judgements. These decisions are based on holding time considerations and client and project specific Data Quality Objectives. The Department Manager, Operations Manager, and/or Quality Assurance Officer may be consulted to evaluate data. Some samples may not be able to be reanalyzed within hold time. In these cases "qualified" data with narration may be advisable after consultation with the client.

In some cases the standard QC requirements listed in this section and in Table 1 may not be sufficient to meet the Data Quality Objectives of the specific project. Much of the work performed at the lab is analyzed in accordance with specific QC requirements spelled out in a project specific Quality Assurance Project Plan (QAPP) or in a program specific Quality Systems Manual (QSM). The reporting limits, acceptance criteria and/or corrective actions may be different than those specified in this SOP. In these cases the appropriate information will be communicated to the Department Manager and/or senior chemists before initiation of the analyses so that specific product codes can be produced for the project. In addition, the work order notes for each project will describe the specific QAPP or QSM to be followed.

#### INITIAL DEMONSTRATION OF PERFORMANCE

8.1 Instrument detection limits (IDL) are determined quarterly for each analyte analyzed on each instrument. This determination requires seven replicate analyses of a reagent blank, performed on three non-consecutive days. The standard deviation of the 21 analyses is multiplied by three to obtain the IDL. For more information on

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performing IDL determinations, refer to the current revision of Katahdin SOP QA-806.

- 8.2 Method detection limits (MDL) are determined annually for each analyte analyzed by each method on each instrument. This determination requires at least seven replicate digestions and analyses of reagent water spiked at 3-5 times the anticipated MDL for each analyte. MDLs differ from IDLs in that the seven replicates are digested prior to analysis, and they may be analyzed on a single day. The standard deviation of the 7 (or more) replicate analyses is multiplied by the Student's t-value to obtain the MDL. For more information on performing MDL determinations, refer to the current revision of Katahdin SOP QA-806.
- 8.3 A Lower Limit of Quantitation Check (LLQC) sample must be prepared and analyzed annually or on an as-needed basis to confirm the laboratory's Practical Quantitation Limits (PQLs). The LLQC sample is equivalent to the PQL standard (Section 8.9) but is carried through the entire sample preparation and analysis process. Element recoveries for the LLQC sample must fall within 70% to 130% of the expected concentrations to confirm the previously established PQLs.

#### ANALYTICAL RUN QC SAMPLES

8.4 Initial instrument calibration: The instrument is calibrated by running a calibration blank and at least three multielement calibration standards. For each element, calibration is performed fitting a single order equation to the calibration data, as follows:

Y=aX + [Blank]

where: Y = Concentration (ug/L)

X = Measured signal intensity (counts per second)

a = Slope of the calibration line

[Blank] = Measured signal intensity of the calibration blank

Fitting the calibration equation through the measured intensity of the calibration blank, rather than through the y-intercept of the line, provides the best calibration accuracy at the low end of the calibration range. When this equation is used, however, the Agilent software does not calculate a calibration coefficient. For this reason, calibration accuracy at the high end of the calibration range is checked by reanalyzing the highest calibration standard as a sample immediately after instrument calibration. Recoveries for all elements must be within 90% to 110% of the true value in the high calibration standard. If the high calibration standard fails, result for the failing elements may not be reported from the run until the problem is corrected and a passing high calibration standard has been analyzed. Calibration accuracy in the middle of the calibration range is verified by analysis of the CCV solution (see Section 8.6 below). Calibration accuracy at the reporting limit is verified by analysis of the PQL Check Standard (see Section 8.7 below).

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- 8.5 The intensities of all internal standards must be monitored for every analysis. When the intensity of any internal standard fails to fall between 70 and 120 percent of the intensity of that internal standard in the initial calibration standard, the following procedure is followed. The sample must be diluted fivefold (1+4) and reanalyzed with the addition of appropriate amounts of internal standards. This procedure must be repeated until the internal standard intensities fall within the prescribed window. The intensity levels of the internal standards for the calibration blanks (ICB and CCBs) and calibration verification standards (ICV and CCVs) must agree within ± 20 percent of the intensity level of the internal standard of the original calibration solution. If they do not agree, terminate the analysis, correct the problem, recalibrate, verify the new calibration, and reanalyze the affected samples.
- 8.6 An Initial Calibration Verification (ICV) solution is analyzed after the initial calibration to check calibration accuracy. The ICV solution is prepared by combining compatible elements from standard sources different than those of the calibration standards and at concentrations within the linear working range of the instrument. The results of the ICV must fall within 90% to 110% of the expected values. If the ICV fails, result for the failing elements may not be reported from the run, unless the ICV recovery is greater than 110% and the sample result is less than the PQL.
- 8.7 Continuing Calibration Verification (CCV) solutions are analyzed after the initial calibration, after every ten samples, and at the end of the analytical run. The CCV solution is prepared using the same standards used for calibration at concentrations near the mid-point of the calibration curve. Results of the CCVs must fall within 90% to 110% of the expected values. If a CCV fails, results for the failing elements in samples bracketed by the failing CCV may not be reported, unless the CCV recovery is greater that 110% and the sample result is less that the PQL. For DoD analyses, results may not be reported without a valid CCV or report flagged results if reanalysis is not possible.
- 8.8 A Practical Quantitation Limit (PQL) Check Standard or low level continuing calibration verification (LLCCV) is analyzed at the beginning of each run (after the ICV and ICB samples) and at the end of each run. Element concentrations in this solution are one-fifth the laboratory's practical quantitation limit (assuming a 5-fold dilution of all digestates during analysis). Element recoveries for the PQL Check Standard must fall within 70% to 130% of the expected values (unless the samples analyzed are for the Department of Defense (80% to 120% recovery limits) or other client-specific limits are imposed). If the PQL Check Standard fails, results for the failing elements may not be reported from the run, unless the PQL Check Standard recovery is greater than the high limit and the sample result is less than the PQL.
- 8.9 A calibration blank solution is analyzed after each ICV and CCV. A calibration blank that is analyzed after the ICV is called an Initial Calibration Blank (ICB). A calibration blank that is analyzed after a CCV is called a Continuing Calibration Blank (CCB). The absolute values of results of ICBs and CCBs must be less than

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the applicable reporting limit (or PQL) for each element. The reporting limit should be determined on a project specific basis, taking into account the data quality objectives for the project. This information must be communicated through a project QAPP and through the Katahdin project manager. When no project specific reporting limit is specified, the laboratory PQL shall be used. Some project specific limits may require reporting down to the MDL or IDL and taking corrective action based on these levels. Results that fall between the PQL and the IDL or MDL must always be flagged as "estimated" with a "J".

- 8.10 If an ICB or a CCB fails the acceptance criteria of less than the reporting limit, results for the failing elements may not be reported from the run until the problem is corrected and a passing ICB or CCB has been analyzed, with the following exception. If the result for an ICB or CCB is greater than the PQL (or reporting limit), sample results that are less than the PQL (or reporting limit) or that are greater than or equal to ten times the measured ICB or CCB concentration may be reported. Also, for failing elements, all samples analyzed after the last passing CCB must be reanalyzed, with the exception noted above.
- 8.11 To obtain analyte data of known quality, it is necessary to measure more than the analytes of interest in order to apply corrections or to determine whether interference corrections are necessary. If the concentrations of interference sources (such as C, Cl, Mo, Zr, W) are such that, at the correction factor, the analyte is less than the limit of quantitation and the concentration of interferents are insignificant, then the data may go uncorrected. Note that monitoring the interference sources does not necessarily require monitoring the interferent itself, but that a molecular species may be monitored to indicate the presence of the interferent. When correction equations are used, all QC criteria must also be met. Extensive QC for interference corrections are required at all times. The monitored masses must include those elements whose hydrogen, oxygen, hydroxyl, chlorine, nitrogen, carbon and sulfur molecular ions could impact the analytes of interest. Interference check solutions ICS-A and ICS-AB are analyzed at the beginning of each run and at least every 12 hours during the run to verify the effectiveness of interference corrections. Solution ICS-A contains high concentrations of interferents (Al, Ca, Fe, Mg, Na, P, K, S, C, Cl, Mo, and Ti) only. These should recover between 80% and 120% of the true value. The measured concentrations of other elements in this solution should be very low, indicating that interfering mass correction equations are adequate. Solution ICS-AB contains interferents at the same high concentrations, and all other analytes at 20 ug/L. Measured recoveries for all analytes should be within 80% to 120% of the true values.

#### PREPARATION BATCH QC SAMPLES

Each digestion batch of twenty or fewer samples will contain a preparation blank and a laboratory control sample. Each batch will also contain one or more of the following QC samples: laboratory control sample duplicate, sample duplicate, matrix spiked sample, or matrix spiked sample duplicate.

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8.12 A preparation blank (PBW or PBS), consisting of reagent water carried through the same process as associated samples, is prepared with each digestion batch of twenty or fewer samples. The results of preparation blanks must be less than the Practical Quantitation Level (PQL) (or project specific reporting limit, if applicable) for each element. If a preparation blank fails, results for the failing elements may not be reported from the digestion batch, and all associated samples must be redigested, with the following exception. If the result for a preparation blank is greater than the PQL or reporting limit, associated sample results that are less than the PQL or reporting limit or that are greater than or equal to ten times the measured preparation blank concentration may be reported.

8.13 A laboratory control sample (LCSW, LCSO, or LCSS), consisting of spiked reagent water or a solid reference material carried through the same process as associated samples, is prepared with each digestion batch of twenty or fewer samples. Results for laboratory control samples must fall within 80% to 120% of the expected value, unless vendor-supplied limits (for solid reference materials) or laboratory-generated statistical limits are available. If a laboratory control sample fails, results for the failing elements may not be reported from the digestion batch, and all associated samples must be redigested, with the following exception. If the recovery of a laboratory control sample is greater than 120%, associated sample results that are less than the PQL or reporting limit may be reported.

#### SAMPLE MATRIX QC SAMPLES

8.14 The relative percent difference (RPD) between matrix duplicate, matrix spike duplicate, or laboratory control duplicate sample results is calculated as follows:

RPD (%) = 
$$\frac{|D_1 - D_2|}{(|D_1 + D_2|)/2} \times 100$$

where:  $D_1$  = First sample or LCS result

D<sub>2</sub>= Second (duplicate) sample or LCS result

A control limit of 20% RPD is applied to duplicate analysis, if the result is greater than 100 times the instrument detection limit. If the matrix duplicate analysis fails, the associated sample result must be flagged on the report of analysis.

8.15 The recovery for each element in a spiked sample must fall within 75% to 125% of the actual value if the result for the unspiked sample is less than four times the amount of spike added. If a recovery fails, the associated sample result must be flagged on the report of analysis. The spike recovery should be calculated as follows:

Recovery (%) = 
$$\frac{S-U}{SA}$$
 \*100%

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where: S = Measured concentration of spiked aliquot

U = Measured concentration of unspiked aliquot

SA = Amount of spike added

8.16 A serial dilution is analyzed to check for chemical or physical interferences. If the analyte concentration of a sample is sufficiently high (minimally, 50 x IDL), the measured concentration of a five-fold dilution (1:5 dilution) of the sample should agree within 90% to 110% of the original determination. The percent difference between the original sample and the serial dilution should be calculated as follows:

Difference (%) = 
$$\frac{|L-S|}{S}$$
 \*100%

where: L = Serial dilution result (corrected for dilution)

S = Original sample result

If the serial dilution analysis fails, a matrix interference should be suspected. The action taken is dependent upon project requirements. The associated sample result may be flagged on the report of analysis, the sample may be reanalyzed at dilution to eliminate the interference, or a post-digestion spike may be performed (see section 8.16).

8.17 An analyte spike that is added to an aliquot of a prepared sample, or its dilution, should be recovered within 80% to 120% of the known value if the result for the unspiked aliquot is less than four times the amount of spike added. If the post-digestion spike is not recovered within these limits, the sample should be diluted and reanalyzed to compensate for the matrix interference or the method standard additions may be employed.

#### 9.0 METHOD PERFORMANCE

The method detection limit (MDL) is defined as the minimum concentration of a substance that can be measured and reported with 99% confidence that the value is above zero. The MDLs shall be determined and verified one time per type of instrument unless otherwise required by the method.

A Limit of Detection (LOD) is an estimate of the minimum amount of a substance that an analytical process can reliably detect. An LOD is analyte and matrix specific and may be laboratory-dependent. LODs must be determined for all parameters for which the laboratory is accredited under the DoD Environmental Laboratory Accreditation Program. LOD's must be verified for every preparation and analytical method combination and on every applicable instrument on a quarterly basis.

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The Limit of Detection (LOQ) is the minimum levels, concentrations, or quantities of a target variable (e.g., target analyte) that can be reported with a specified degree of confidence. The LOQ shall be set at the lowest point in the calibration curve for all analyses utilizing an initial calibration. LOQ's must be verified quarterly for every preparation and analytical method combination and on every applicable instrument on a quarterly basis for all parameters included in the DoD Scope of Accreditation. The LOQ must be verified at least once annually if the analysis is not included in the DoD Scope of Accreditation.

MDLs are filed with the Organic Department Manager and then with the QAO. LOD and LOQ verifications are filed with the QAO

Refer to the current revision of Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, for procedures on determining the MDL.

Refer to the current revisions of USEPA Method 6020 for other method performance parameters and requirements.

#### 10.0 APPLICABLE DOCUMENTS/REFERENCES

Test Methods for Evaluating Solid Waste, Physical/Chemical Methods, USEPA SW846, 3<sup>rd</sup> Edition, Final Updates I, II, IIA, IIB, III, IIIA, IIIB and IV, February 2007, Method 6020A.

The NELAC Institute, Laboratory Accreditation Standards, Volume 1, Management and Technical Requirements for Laboratories Performing Environmental Analysis, 10/06/2010.

Department of Defense Quality Systems Manual for Environmental Laboratories (DOD QSM), Current Version.

Katahdin SOP CA-101, Equipment Maintenance and Troubleshooting, current revision.

Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, current revision

Agilent 7500 ICP-MS ChemStation Operator's Manual, Agilent Technologies, Inc., 2000.

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## TABLE 1

## QC REQUIREMENTS

QC Sample	Minimum Frequency	Acceptance Criteria	Corrective Action
Initial Calibration, minimum 3 points plus a calibration blank.	Daily prior to sample analysis.	Correlation coefficient (r) ≥ 0.998	Recalibrate
Initial Calibration Verification (ICV), prepared from a second source.	Before beginning a sample run.	Recovery within <u>+</u> 10% of true value.	Do not use results for failing elements, unless ICV >110% and sample result < PQL/reporting limit.
Initial Calibration Blank (ICB)	Immediately after the ICV.	Absolute value of ICB < PQL or project specific reporting limit.	Do not use results if sample > PQL/reporting limit and < 10x ICB level.
PQL Standard or LLCCV	At beginning and end of run	70-130% of true value	Do not use results for failing elements, unless PQL rec.> upper limit and sample result < PQL/reporting limit.
Continuing Calibration Verification (CCV)	At beginning of run, after every 10 samples, and at end of run.	Recovery within ± 10% of true value.	1) Do not use bracketed sample results for failing elements, unless CCV >110% and sample result < PQL/reporting limit. 2) Investigate and correct problem.
Continuing Calibration Blank (CCB)	Immediately after every CCV	Absolute value of CCB < PQL or project specific reporting limit.	Do not use sample results if ≥ PQL/reporting limit and < 10x CCB level.
Interference Check Solution A (ICS-A)	Before analyzing samples, and every 12 hours during a run.	Interferents: Recovery within ± 20% of true value. Analytes: No criteria established (Project specific criteria may apply)	Do not use sample results for failing elements.
Interference Check Solution AB (ICS-AB)	Before analyzing samples, and every 12 hours during a run.	Recovery within ± 20% of true value.	Do not use sample results for failing elements, unless ICSAB >120% and sample result < PQL/reporting limit.
Preparation Blank (PBW/PBS)	One per digestion batch of 20 or fewer samples.	Less than PQL (standard practice), or based on the project specific guidelines.	<ol> <li>Investigate source of contamination.</li> <li>Redigest and reanalyze all associated samples if sample concentration ≥ PQL and &lt;10x the blank conc.</li> </ol>
Laboratory Control Sample (LCSW/LCSS/LCSO)	At least one per digestion batch of 20 or fewer samples.	Recovery within ± 20% of true value, unless vendor-supplied or statistical limits have been established.	Investigate source of problem.     Redigest and reanalyze all associated samples, unless LCS >120% and sample result < PQL.

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## TABLE 1 (continued)

## QC REQUIREMENTS

QC Sample	Minimum Frequency	Acceptance Criteria	Corrective Action
Duplicate Sample (D), Matrix Spike Duplicate (P), or LCS Duplicate (LC2W/LC2S/LC2O)	See section 8.11	1) RPD ≤ 20%, if sample > 100x IDL.	Flag results
Post-Digestion Matrix Spike (A)	When serial dilution fails and analyte concentration < 100 x MDL.	Recovery <u>+</u> 20% of true value, if sample < 4x spike added.	Flag results and/or analyze sample by method of standard additions.
Serial Dilution (L)	1 per digestion batch	If original sample result is at least 50x IDL, 5-fold dilution must agree within ± 10% of the original result.	Flag result or dilute and reanalyze sample to eliminate interference.
Internal Standard (IS)	Appropriate IS required for all analytes in all samples. Mass of IS must be <50 amu different from that of analyte.	1) For each sample, IS intensity within 70%-120% of that of initial calib. blank. 2) For ICV, ICB, CCV, and CCB, IS intensity within 80%-120% of that in initial calib. blank.	Do not use results for failing elements.
Instrument Detection Limit (IDL) Study	Quarterly.	IDL < MDL PQL at least 2-3x IDL	Repeat IDL study.     Raise PQL.
Method Detection Limit (MDL) Study		A-806, "Method Detection Lines and Verifications", current	nit, Instrument Detection Limit and revision.
Lower Limit of Quantitation Check (LLQC) Sample	Digest and analyze annually or as needed to confirm PQLs	70% - 130% of true value	Reevaluate PQLs
Method of Standard Additions	When matrix interference is suspected	r <u>&gt; 0</u> .995	Dilute and reanalyze sample to eliminate interference.

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## TABLE 2

## DoD QSM QC REQUIREMENTS

QC Check	Minimum	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
QO OHOOK	Frequency	71000ptaniou Ontonia	001100111071011011	i lagging official	
Demonstrate acceptable analytical capability	Prior to using any test method and at any time there is a change in instrument type, personnel, test method, or sample matrix.	QC acceptance criteria published by DoD, if available; otherwise method-specified criteria.	Recalculate results; locate and fix problem, then rerun demonstration for those analytes that did not meet criteria.	NA.	This is a demonstration of analytical ability to generate acceptable precision and bias per the procedure in Appendix C. No analysis shall be allowed by analyst until successful demonstration of capability is complete.
LOD determination and verification LOQ	Refer to current revision of SOP QA-806 Refer to current				
establishment and verification	revision of SOP QA-806				
Instrument detection limit (IDL) study	At initial set-up and after significant change in instrument type, personnel, test method, or sample matrix.	IDLs shall be ≤ LOD.	NA.	NA.	Samples may not be analyzed without a valid IDL.
Tuning	Prior to ICAL.	Mass calibration ≤ 0.1 amu from the true value; Resolution < 0.9 amu full width at 10% peak height; For stability, RSD ≤ 5% for at least four replicate analyses.	Retune instrument then reanalyze tuning solutions.	Flagging criteria are not appropriate.	No analysis shall be performed without a valid MS tune.
Initial calibration (ICAL) for all analytes (minimum one high standard and a calibration blank)	Daily ICAL prior to sample analysis.	If more than one calibration standard is used, r ≥ 0.995.	Correct problem, then repeat ICAL.	Flagging criteria are not appropriate.	Problem must be corrected. No samples may be run until ICAL has passed.
Second source calibration verification	Once after each ICAL, prior to beginning a sample run.	Value of second source for all analytes within ± 10% of true value.	Verify second source standard. Rerun second source verification. If that fails, correct problem and repeat ICAL.	Flagging criteria are not appropriate.	Problem must be corrected. No samples may be run until calibration has been verified.

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## TABLE 2 (cont)

## DoD QSM QC REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Continuing calibration verification (CCV)	After every 10 field samples and at the end of the analysis sequence.	All analytes within ± 10% of true value.	Correct problem, rerun calibration verification. If that fails, then repeat ICAL. Reanalyze all samples since the last successful calibration verification.	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply Q-flag to all results for the specific analyte(s) in all samples since the last acceptable calibration verification.	Problem must be corrected. Results may not be reported without a valid CCV. Flagging is only appropriate in cases where the samples cannot be reanalyzed.
Low-level calibration check standard	Daily, after one-point ICAL.	Within ± 20% of true value.	Correct problem, then reanalyze.	Flagging criteria are not appropriate.	No samples may be analyzed without a valid low-level calibration check standard. Low-level calibration check standard should be less than or equal to the reporting limit.
Linear dynamic range or high- level check standard	Every 6 months.	Within ±10% of true value.	NA.	NA.	
Method blank	One per preparatory batch.	No analytes detected > ½ RL (> RL for common lab contaminants) and > 1/10 the amount measured in any sample or 1/10 the regulatory limit (whichever is greater). For negative blanks, absolute value < LOD. Blank result must not otherwise affect sample results.	Correct the problem. Report sample results that are <lod or="">10x the blank concentration. Reprepare and reanalyze the method blank and all associated samples with results &gt; LOD and &lt; 10x the contaminated blank result.</lod>	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply B-flag to all results for the specific analyte(s) in all samples in the associated preparatory batch.	Problem must be corrected. Results may not be reported without a valid method blank. Flagging is only appropriate in cases where the samples cannot be reanalyzed.
Calibration blank	Before beginning a sample run, after every 10 samples, and at end of the analysis sequence.	No analytes detected > LOD. For negative blanks, absolute value < LOD.	Correct problem. Reprep and reanalyze calibration blank. All samples following the last acceptable calibration blank must be reanalyzed.	Apply B-flag to all results for specific analyte(s) in all samples associated with the blank.	
Interference check solutions (ICS-A and ICS-AB)	At the beginning of an analytical run and every 12 hours.	ICS-A: Absolute value of concentration for all non-spiked analytes < LOD (unless they are a verified trace impurity from one of the spiked analytes); ICS-AB: Within ± 20% of true value.  May use < LOD for some projects.	Terminate analysis, locate and correct problem, reanalyze ICS, reanalyze all samples.	If corrective action fails, apply Q-flag to all results for specific analyte(s) in all samples associated with the ICS.	

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## TABLE 2 (cont)

## DoD QSM REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
LCS containing all analytes to be reported	One per preparatory batch.	Water: Recovery must be within + 20% of the true value Soil: Recovery must be within vendor supplied limits (varies by lot).	Correct problem, then reprep and reanalyze the LCS and all samples in the associated preparatory batch for failed analytes, if sufficient sample material is available.	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply Q-flag to specific analyte(s) in all samples in the associated preparatory batch.	Problem must be corrected. Results may not be reported without a valid LCS. Flagging is only appropriate in cases where the samples cannot be reanalyzed.
Matrix spike (MS)	One per preparatory batch per matrix	For matrix evaluation, use recovery must be within + 20% of the true value.	Examine the project- specific DQOs. If the matrix spike falls outside of DoD criteria, additional quality control tests (dilution test and post-digestion spike addition) are required to evaluate matrix effects.	For the specific analyte(s) in the parent sample, apply J-flag if acceptance criteria are not met.	For matrix evaluation only. If MS results are outside the LCS limits, the data shall be evaluated to determine the source of difference and to determine if there is a matrix effect or analytical error.
Matrix spike duplicate (MSD) or sample duplicate	One per preparatory batch per matrix.	MSD: For matrix evaluation use recovery must be within + 20% of the true value. MSD or sample duplicate: RPD < 20% (between MS and MSD or sample and sample duplicate).	Examine the project- specific DQOs. Contact the client as to additional measures to be taken.	For the specific analyte(s) in the parent sample, apply J-flag if acceptance criteria are not met.	The data shall be evaluated to determine the source of difference.
Dilution test	One per preparatory batch.	For samples with concentrations > 50 x LOQ then five-fold dilution must agree within ± 10% of the original measurement.	Perform post-digestion spike addition.	Flagging criteria are not appropriate.	Only applicable for samples with concentrations > 50 x LOQ.
Post digestion spike addition	When dilution test fails or analyte concentration for all samples < 50 x LOD.	Recovery within 75-125%	Run all associated samples in the preparatory batch by method of standard additions (MSA) or see flagging criteria.	For the specific analyte(s) in the parent sample, apply J-flag if acceptance criteria are not met.	Spike addition should produce a concentration of 10 – 100 x LOQ.
Method of standard additions (MSA)	When matrix interference is confirmed.	NA.	NA.	NA.	Document use of MSA in the case narrative.
Internal standards (IS)	Every sample.	IS intensity within 30- 120% of intensity of the IS in the ICAL.	Flagging criteria are not appropriate.	Reanalyze sample at 5- fold dilution with addition of appropriate amounts of internal standards.	
Results reported between DL and LOQ	NA.	NA.	NA.	Apply J-flag to all results between DL and LOQ.	

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# TABLE 3 SUMMARY OF METHOD MODIFICATIONS

TOPIC	KATAHDIN SOP CA-627-07	METHOD 6020, current revision
Apparatus/Materials		
Reagents		
Sample preservation/ handling		
Procedures		
QC - Spikes		
QC - LCS		
QC - Accuracy/Precision		
QC - MDL		
QC - Calibration Blanks	Acceptance criteria employed for 6020: ± PQL	Acceptance criteria stated in 6020: <10% of PQL

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TABLE 4
ISOTOPES MONITORED AND CORRECTION EQUATIONS USED FOR USEPA METHOD 6020

Element Class	Element	Sym- bol	Isotopes Monitored	Correction Equations [See note 1]
	Aluminum	Al	27	
	Antimony	Sb	121, 123	
	Arsenic	As	75	$^{75}$ As = $(75)$ *1 - $(77)$ *2.95 + $(82)$ *2.548 - $(83)$ *2.571
				[See note 2]
	Barium	Ва	135, 137	
	Beryllium	Be	9	
	Boron	В	11	
	Cadmium	Cd	106, 108, 111,	<sup>111</sup> Cd = (111)*1 – (108)*1.073 + (106)*0.764 [See
			114	note 3]
				<sup>114</sup> Cd = (114)*1 – (118)*0.0268 [See note 4]
	Calcium	Ca	44	<sup>44</sup> Ca = (44)*1 – (88)*0.0169 [See note 7]
	Chromium	Cr	52, 53	
	Cobalt	Со	59	
	Copper	Cu	63, 65	- E4
	Iron	Fe	54, 56, 57	<sup>54</sup> Fe = (54)*1 – (52)*0.0282 [See note 8]
				<sup>57</sup> Fe = (57)*1 – (43)*0.03 [See note 9]
Analytes	Lead	Pb	206, 207, 208	<sup>208</sup> Pb = (208)*1 + (206)*1 + (207)*1 [See note 5]
	Magnesium	Mg	25	
	Manganese	Mn	55	00
	Molybdenum	Мо	98	<sup>98</sup> Mo = (98)*1 – (99)*0.146 [See note 10]
	Nickel	Ni	60, 61	
	Potassium	K	39	92 _
	Selenium	Se	82	<sup>82</sup> Se = (82)*1 – (83)*1.009 [See note 11]
	Silver	Ag	107, 109	
	Sodium	Na	23	
	Strontium	Sr	88	
	Thallium	TI	203, 205	
	Thorium	Th	232	
	Tin	Sn	118, 120	
	Tungsten	W	182	
	Uranium	U	238	51.
	Vanadium	V	51	$^{51}V = (51)*1 - (53)*2.95 + (52)*0.333$ [See note 12]
	Zinc	Zn	66, 67, 68	
	Bismuth	Bi	209	
	Germanium	Ge	72	115.
Internal	Indium	In	115	<sup>115</sup> In = (115)*1 – (118)*0.016 [See note 6]
Stan-	Lithium	Li	6	
dards.	Scandium	Sc	45	
	Terbium	Tb	159	
	Yttrium	Υ	89	

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TABLE 4 (continued)

#### ISOTOPES MONITORED AND CORRECTION EQUATIONS USED FOR USEPA METHOD 6020

#### Notes:

- 1) Numbers in parentheses, e.g "(51)", indicate measured counts at the indicated mass.
- 2) Corrects for ArCl interference, taking into account secondary interferences from Se and Kr
- 3) Corrects for MoO interference, taking into account secondary interference from <sup>108</sup>Cd
- 4) Corrects for Sn interference
- 5) Corrects for variations in isotopic composition of lead
- 6) Corrects for Sn interference
- 7) Corrects for interference from <sup>88</sup>Sr<sup>2+</sup>
- 8) Corrects for Cr interference
- 9) Corrects for Ca interference
- 10) Corrects for Ru interference
- 11) Corrects for Kr interference
- 12) Corrects for CIO, taking into account secondary interference from Cr

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TABLE 5

PREPARATION OF CALIBRATION AND QUALITY CONTROL STANDARDS

Sample or Solution Name	Component Solution Name	Source of Component	Amount of Component Added per 100 mL Final Volume (mL)
Calibration Standard 1	CL-CAL-3	Spex Industries	0.005
<b>(S1)</b> (1.0% HNO <sub>3</sub> /	ICP-MS-MIX-Z	Lab Prepared	0.01
0.5% HCI)	ICP-MS CAL 1	Lab Prepared	0.025
Calibration Standard 2	CL-CAL-3	Spex Industries	0.05
<b>(S2)</b> (1.0% HNO <sub>3</sub> /	ICP-MS-MIX-Z	Lab Prepared	0.10
0.5% HCI)	ICP-MS CAL 1	Lab Prepared	0.25
Calibration Standard 3	CL-CAL-3	Spex Industries	0.25
(S3) / CCV (1.0% HNO <sub>3</sub> /	ICP-MS-MIX-Z	Lab Prepared	0.50
0.5% HCI)	ICP-MS CAL 1	Lab Prepared	1.25
Calibration Standard 4 (S4) / High Standard	CL-CAL-3	Spex Industries	0.50
` (1.0% HNO₃ /	ICP-MS-MIX-Z	Lab Prepared	1.0
0.5% HCI)	ICP-MS CAL 1	Lab Prepared	2.5
	CL-ICS-1,CL-ICS-4, CL-ICS-5	Spex Industries	0.20 of each
Initial Calibration	CL-ICS-3	Spex Industries	2.0
Verification (ICV) (1.0% HNO <sub>3</sub> /	1000 mg/L Si	Inorganic Ventures	0.040
0.5% HCI)	1000 mg/L Al	Inorganic Ventures	0.038
	1000 mg/L B, W	Inorganic Ventures	0.002 of each
Continuing Calibration	CL-CAL-3	Spex Industries	0.25
Verification (CCV) (1.0% HNO3/ 0.5% HCI)	ICP-MS-MIX-Z, ICP-MS CAL 1, ICP-MS-MIX-Y	Lab Prepared	0.50 of each

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#### TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020

## TABLE 5 (continued)

## PREPARATION OF CALIBRATION AND QUALITY CONTROL STANDARDS

Sample or Solution Name	Component Solution Name	Source of Component	Amount of Component Added per 100 mL Final Volume (mL)
Practical Quantitation Limit Solution (PQL) (1.0% HNO <sub>3</sub> / 0.5% HCl)	ICP-MS PQL Intermediate	Lab Prepared	0.1
Interference Check Solution A (ICS-A) (1.0% HNO <sub>3</sub> / 0.5% HCI)	6020ICS-0A	Inorganic Ventures	10.0
Interference Check	6020ICS-0A	Inorganic Ventures	10.0
Solution AB (ICS-AB)	ICP-MS-CAL 1	Lab Prepared	1.0
(1.0% HNO <sub>3</sub> / 0.5% HCl)	ICP-MS ICSAB Intermediate	Lab Prepared	1.0
P/A Tuning Solution (1.0% HNO <sub>3</sub> /	1000 mg/L Co, Cr, Mo, Mn, Pb, Sb, Sr, U, V	High Purity Standards	0.02
0.5% HCI)	10,000 mg/L AI, K, Na	High Purity Standards	0.002
Instrument Tuning Solution (1.0% HNO <sub>3</sub> / 0.5% HCl)	ICP-MS-TS-2	High Purity Standards	0.10
Internal Standard Solution (5.0% HNO <sub>3</sub> / 0.5% HCl)	Internal Standard Mix	Spex Industries	10
Method Tuning Solution	ICP-MS Method Tune Intermediate	Lab Prepared	1.0
(1.0% HNO₃ / 0.5% HCl)	Internal Standard Mix 1	Spex Industries	1.0

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# TABLE 6 PREPARATION OF INTERMEDIATE STANDARDS

Sample or Solution Name	Component Solution Name	Source of Component	Amount of Component Added per 100 mL Final Volume (mL)
	10,000 mg/L K, Na	High Purity Standards or Inorganic Ventures	2.0 of each
	10,000 mg/L Si	High Purity Standards or Inorganic Ventures	1.0
	10,000 mg/L Al	High Purity Standards or Inorganic Ventures	0.60
ICP-MS PQL Intermediate	1000 mg/L B	High Purity Standards or Inorganic Ventures	0.40
(1.0% HNO <sub>3</sub> / 0.5% HCl)	10,000 mg/L Ca, Fe, Mg 1000 mg/L Zn	High Purity Standards	0.20 of each
	1000 mg/L As, Se, V, W, Sr, Sn, Mo, Cr	High Purity Standards or Inorganic Ventures	0.10 of each
	1000 mg/L Cu	High Purity Standards	0.06
	1000 mg/L Ba, Mn, Ni	High Purity Standards	0.04 of each
	1000 mg/L U, Be, Cd, Co, Ag, Tl, Pb, Sb	High Purity Standards	0.02 of each
ICP-MS CAL 1 (1.0% HNO <sub>3</sub> / 0.5% HCI)	1000 mg/L Ag, As, Ba, Be, Cd, Co, Cr, Cu, Mn, Mo, Ni, Pb, Sb, Se, Tl, U, V, Zn	High Purity Standards	0.2 of each
0.5% HOI)	10,000 mg/L AI	High Purity Standards or Inorganic Ventures	0.02
	10,000 mg/L K, Na, Fe, Mg, Ca	High Purity Standards or Inorganic Ventures	5.0 of each
ICP-MS-MIX-Z (1.0% HNO <sub>3</sub> /	10,000 mg/L Si	High Purity Standards or Inorganic Ventures	1.0
0.5% HCI)	10,000 mg/L AI	High Purity Standards or Inorganic Ventures	0.95
	1000 mg/L B, Sn, Sr, W	High Purity Standards or Inorganic Ventures	0.50 of each

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## TABLE 6 (Cont'd)

## PREPARATION OF INTERMEDIATE STANDARDS

ICP-MS-MIX-Y	10,000 mg/L Al	High Purity Standards or Inorganic Ventures	0.030
(1.0% HNO3/ 0.5% HCI)	1000 mg/L As, Ba, Cr, Cu, Mn, Mo, Ni, Pb, Se, Sb, V, Zn	High Purity Standards or Inorganic Ventures	0.30 of each
ICP-MS ICSAB Intermediate	10,000 mg/L Si	High Purity	0.50
(1.0% HNO <sub>3</sub> / 0.5% HCI)	1,000 mg/L B, Sn, Sr, W	High Purity or Inorganic Ventures	0.20 each
ICP-MS Method Tune Intermediate (1.0% HNO <sub>3</sub> / 0.5% HCl)	1000 mg/L Be, Co, Pb, Tl 10,000 mg/L Mg	High Purity Standards or Inorganic Ventures	0.1 of each

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# TABLE 7 ELEMENT CONCENTRATIONS IN WORKING STANDARDS

			CONC	CENTRATION	IN SOLUTIO	N, ug/L		
Element	<b>S</b> 1	S2	S3	S4	ICV	PQL	CCV	P/A Tune Soln.
Aluminum	10.0	100.0	500.0	1000.0	400.0	60.0	500.0	200
Antimony	0.5	5.0	25.0	50.0	20.0	0.2	25.0	200
Arsenic	0.5	5.0	25.0	50.0	20.0	1.0	25.0	
Barium	0.5	5.0	25.0	50.0	20.0	0.4	25.0	
Beryllium	0.5	5.0	25.0	50.0	20.0	0.2	10.0	
Boron	0.5	5.0	25.0	50.0	20.0	4.0	25.0	
Cadmium	0.5	5.0	25.0	50.0	20.0	0.2	10.0	
Calcium	100.0	1000.0	5000.0	10000.0	4000.0	20.0	5000.0	
Chromium	0.5	5.0	25.0	50.0	20.0	1.0	25.0	200
Cobalt	0.5	5.0	25.0	50.0	20.0	0.2	10.0	200
Copper	0.5	5.0	25.0	50.0	20.0	0.6	25.0	
Iron	100.0	1000.0	5000.0	10000.0	4000.0	20.0	5000.0	
Lead	0.5	5.0	25.0	50.0	20.0	0.2	25.0	200
Magnesium	100.0	1000.0	5000.0	10000.0	4000.0	20.0	5000.0	
Manganese	0.5	5.0	25.0	50.0	20.0	0.4	25.0	200
Molybdenum	0.5	5.0	25.0	50.0	40.0	1.0	25.0	200
Nickel	0.5	5.0	25.0	50.0	20.0	0.4	25.0	
Potassium	100.0	1000.0	5000.0	10000.0	4000.0	200.0	5000.0	200
Selenium	0.5	5.0	25.0	50.0	20.0	1.0	25.0	
Silicon	10.0	100.0	500.0	1000.0	400.0	100.0	500.0	
Silver	0.5	5.0	25.0	50.0	20.0	0.2	10.0	
Sodium	100.0	1000.0	5000.0	10000.0	4000.0	200.0	5000.0	200
Strontium	0.5	5.0	25.0	50.0	20.0	1.0	25.0	200
Thallium	0.5	5.0	25.0	50.0	20.0	0.2	10.0	
Tin	0.5	5.0	25.0	50.0	20.0	1.0	25.0	
Tungsten	0.5	5.0	25.0	50.0	20.0	1.0	25.0	
Uranium	0.5	5.0	25.0	50.0	20.0	0.2	10.0	200
Vanadium	0.5	5.0	25.0	50.0	20.0	1.0	25.0	200
Zinc	0.5	5.0	25.0	50.0	20.0	2.0	25.0	

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#### TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020

## TABLE 7 (continued)

## ELEMENT CONCENTRATIONS IN WORKING STANDARDS

	CONCENTRATION IN SOLUTION, ug/L				
Element	ICSA <sup>1</sup>	ICSAB <sup>1</sup>	Internal Std Solution	Method Tune Solution	Instrument Tuning Solution
Aluminum	100000	100000			
Antimony		20			
Arsenic		20			
Barium		20		10	
Beryllium		20			
Boron		20			
Cadmium		20			
Calcium	100000	100000			
Chromium		20			
Cobalt		20		10	
Copper		20			
Iron	100000	100000			
Lead		20		10	
Magnesium	100000	100000		100	
Manganese		20			
Molybdenum	2000	2000			
Nickel		20			
Potassium	100000	100000			
Selenium		20			
Silver		20			
Sodium	100000	100000			
Strontium		20			
Thallium		20		10	10.0
Tin		20			
Tungsten		20			
Uranium		20			
Vanadium		20			
Zinc		20			
Bismuth			1000.0	10	
Germanium			1000.0	10	
Indium				10	
Lithium ( <sup>6</sup> Li)			1000.0	10	
Scandium			1000.0	10	
Terbium			1000.0	10	
Yttrium			1000.0	10	10.0
Cerium					10.0

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TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020

TABLE 7 (continued)

## **ELEMENT CONCENTRATIONS IN WORKING STANDARDS**

	CONCENTRATION IN SOLUTION, ug/L				
Element	ICSA <sup>1</sup>	ICSAB <sup>1</sup>	Internal Std Solution	Method Tune Solution	Instrument Tuning Solution
Lithium					10.0

<sup>1)</sup> Solution also contains 1000 mg/L Chloride, 200 mg/L Carbon, and 100 mg/L Phosphorus and Sulfur, and 2mg/L Titanium.

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# TABLE 8 ELEMENT CONCENTRATIONS IN INTERMEDIATE STANDARDS

	CONCENTRATION IN SOLUTION, mg/L						
ELEMENT	MS-MIX-Z	ICP-MS PQL Intermediate	ICP-MS-MIX-Y	ICP-MS Method Tune Intermediate	ICP-MS CAL 1	ICP-MS ICSAB Intermediate	
Aluminum	95.0	6.0	3.0		0.2		
Antimony		0.02	3.0		0.2		
Arsenic		0.10	3.0		0.2		
Barium		0.04	3.0		0.2		
Beryllium		0.02		1.0	0.2		
Boron	5.0	4.0				0.2	
Cadmium		0.02			0.2		
Calcium	500	2.0					
Chromium		0.10	3.0		0.2		
Cobalt		0.02		1.0	0.2		
Copper		0.06	3.0		0.2		
Iron	500	2.0					
Lead		0.02	3.0	1.0	0.2		
Magnesium	500	2.0		10.0			
Manganese		0.04	3.0		0.2		
Molybdenum		0.10	3.0		0.2		
Nickel		0.04	3.0		0.2		
Potassium	500	20.0					
Selenium		0.10	3.0		0.2		
Silicon	100	10.0				5.0	
Silver		0.02			0.2		
Sodium	500	20.0					
Strontium	5.0	0.10				0.2	
Thallium		0.02		1.0	0.2		
Tin	5.0	0.10				0.2	
Tungsten	5.0	0.10				0.2	
Uranium		0.020			0.2		
Vanadium		0.10	3.0		0.2		
Zinc		0.20	3.0		0.2		

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TABLE 9
ELEMENT CONCENTRATIONS IN STOCK STANDARDS

	CONCENTRATION IN SOLUTION, mg/L					
Element	Instrument Calibration Standard 3 (Spex)	CL-ICS-1 (Spex)	CL-ICS-3 (Spex)	CL-ICS-4 (Spex)	CL-ICS-5 (Spex)	
Aluminum		10.0				
Antimony		10.0				
Arsenic		10.0				
Barium		10.0				
Beryllium		10.0				
Boron						
Cadmium		10.0				
Calcium	1000		200.0			
Chromium		10.0				
Cobalt		10.0				
Copper		10.0				
Iron	1000		200.0			
Lead		10.0				
Magnesium	1000		200.0			
Manganese		10.0				
Molybdenum				10.0	10.0	
Nickel		10.0				
Potassium	1000		200.0			
Selenium		10.0				
Silver		10.0				
Sodium	1000		200.0			
Strontium					10.0	
Thallium		10.0				
Thorium				10.0		
Tin					10.0	
Tungsten						
Uranium				10.0		
Vanadium		10.0				
Zinc		10.0				

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TITLE: TRACE METALS ANALYSIS BY ICP-MS USING USEPA METHOD 6020

## TABLE 9 (continued)

## **ELEMENT CONCENTRATIONS IN STOCK STANDARDS**

CONCENTRATION IN SOLUTION, ug/L			ug/L
Element	6020ICS-0A <sup>1</sup> (Inorganic Ventures)	Internal Standard Mix 1 (Spex)	ICP-MS-TS-2 (High Purity)
Aluminum	1000		
Arsenic			
Cadmium			
Calcium	1000		
Chromium			
Cobalt			
Copper			
Iron	1000		
Magnesium	1000		
Manganese			
Molybdenum	20.0		
Nickel			
Potassium	1000		
Silver			
Sodium	1000		
Zinc			
Bismuth		1000	
Cerium			10000
Germanium		1000	
Indium		1000	
Lithium			10000
Lithium ( <sup>6</sup> Li)		1000	
Scandium		1000	
Terbium		1000	
Thallium			10000
Yttrium		1000	10000

<sup>1)</sup> Solution also contains 10000 mg/L Chloride, 2000 mg/L Carbon, and 1000 mg/L Phosphorus and Sulfur, and 20 mg/L Titanium.

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## TABLE 10

## REQUIRED ANALYTICAL SEQUENCE

Sequence Number	Standard/Sample	Purpose
1	Method Tuning Solution	Verify mass calibration and resolution
2	S0 (Calibration Blank)	Initial calibration
3	S1 (Calibration Standard)	Initial calibration
4	S2 (Calibration Standard)	Initial calibration
5	S3 (Calibration Standard)	Initial calibration
6	S4 (Calibration Standard)	Initial calibration
7	ICV (Initial Calibration Verification)	Check calibration accuracy
8	ICB (Initial Calibration Blank)	Check calibration accuracy
9	PQL (Practical Quantitation Limit)	Check calibration accuracy at low concentration
10	ICS-A (Interference Check Solution A)	Verify accuracy of mass correction equations
11	ICS-AB (Interference Check Solution AB)	Verify accuracy of mass correction equations
12	CCV (Continuing Calibration Verification)	Check calibration stability
13	CCB (Continuing Calibration Blank)	Check calibration stability
14-23	Analyze up to 10 samples	
24	CCV (Continuing Calibration Verification)	Check calibration stability
25	CCB (Continuing Calibration Blank)	Check calibration stability
	Continue analyzing sequences of up to 10 samples, followed by a CCV and a CCB  After last analytical sample, analyze PQL, followed by a CCV and a CCB	

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# TABLE 11 INSTRUMENT OPERATING CONDITIONS

	Acquisition Mode	Spectrum	
	Points per Mass	3	
	Number of Replicates	3	
	Detector Mode	Auto for all elements	
		0.10 sec for Li, B, <sup>29</sup> Si, Sc, V, Cr, Mn, Ni, Cu, Zn, Y, Mo, Ag, In, Sn, Sb, Ba, Tb, W, Tl, Pb, Bi, Th, U	
Data Acquisition Program	Integration Time per Point (for	0.30 sec for Be, As, Cd, Ge	
	listed masses and their correction	0.010 sec for Na, Al, K, <sup>28</sup> Si	
	masses)	0.030 for Ca, Fe, Sr	
		1.00 sec for Se	
		0.050 sec for Mg, Co	
	Spray Chamber Temperature	2° C	
	Total Acquisition Time	105 sec for 3 replicates	
Peristaltic Pump Program	Analysis Speed	0.15 rps	
	Uptake Speed	0.15 rps	
Before Acquisition	Uptake Time	5 sec	
	Stabilization Time	15 sec	
	Rinse Speed	0.15 rps	
After Acquisition (Probe Rinse)	Rinse Time (sample)	5 sec	
	Rinse Time (standard)	5 sec	
	Rinse Vial	1	
After Acquisition (Rinse)	Uptake Speed	0	
. ,	Uptake Time	0 sec	
	Stabilization Time	0 sec	
	All quantitation masses	Y=ax+(blank)	
Calibration Curve fit	All internal standard masses	(Excluded)	
	All interference correction masses	(Excluded)	
Reporting Parameters	QC Reports	On-Printer	
1136013	All Other Reports	Off	

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## TABLE 12 INSTRUMENT TUNE SPECIFICATIONS

	1: > F000 ata/0 4 a a a/40 mmb		
	Li >5000 cts/0.1 sec/10 ppb		
Sensitivity	Y >10,000 cts/0.1 sec/10 ppb		
	TI >5000 cts/0.1 sec/10 ppb		
	Li <8% RSD (0.1 sec integration time)		
Precision	Y <5% RSD (0.1 sec integration time)		
	TI <5% RSD (0.1 sec integration time)		
Oxides	<1.0%		
Doubly Charged (Ce <sup>++</sup> /Ce <sup>+</sup> )	<2.0%		
	Li <15 cps		
Background	Y <15 cps		
	TI <15 cps		
Mass Resolution	Width at 10% peak height: 0.7-0.8 amu		
	Li ±0.1 amu of nominal mass		
Mass Axis	Y ±0.1 amu of nominal mass		
	TI ±0.1 amu of nominal mass		

TABLE 13
METHOD TUNE SPECIFICATIONS

Precision	≤5% RSD of 4 replicates	
Mass Resolution	Width at 10% peak height: <0.9 amu	
Mass Calibration	±0.1 amu of nominal mass	

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TABLE 14

REPORTED ISOTOPES AND INTERNAL STANDARDS

ELEMENT	MASS	INTERNAL STANDARD (mass)
Aluminum	27	Scandium (45)
Antimony	123	Terbium (159)
Arsenic	75	Germanium (72)
Barium	135	Terbium (159)
Beryllium	9	Lithium (6)
Boron	11	Lithium (6)
Cadmium	114	Germanium (72)
Calcium	44	Scandium (45)
Chromium	52	Germanium (72)
Cobalt	59	Germanium (72)
Copper	65	Germanium (72)
Iron	57	Germanium (72)
Lead	208	Bismuth (209)
Magnesium	25	Scandium (45)
Manganese	55	Germanium (72)
Molybdenum	98	Germanium (72)
Nickel	60	Germanium (72)
Potassium	39	Scandium (45)
Selenium	82	Germanium (72)
Silicon	29	Scandium (45)
Silver	107	Germanium (72)
Sodium	23	Scandium (45)
Strontium	88	Germanium (72)
Thallium	203	Bismuth (209)
Thorium	232	Bismuth (209)
Tin	118	Terbium (159)
Tungsten	182	Terbium (159)
Uranium	238	Bismuth (209)
Vanadium	51	Germanium (72)
Zinc	66	Germanium (72)

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#### ATTACHMENT 1

#### HARDNESS BY CALCULATION

As referenced in "Standard Methods for the Examination if Water and Wastewater," Methods 2340 A & B, Hardness Introduction and Hardness by Calculation, American Public Health Association, 18<sup>th</sup> Edition, Revised 1992, total hardness is the sum of the calcium and magnesium concentrations, both expressed as calcium carbonate, in milligrams per liter.

Once the calcium and magnesium concentrations have been determined by EPA methods 6010, 6020, 200.7 or 200.8, the total hardness of an aqueous sample may be calculated as follows:

Total Hardness, mg equivalent  $CaCO_3/L = 2.497$  (Ca, mg/L) + 4.118 (Mg, mg/L)

The calcium hardness of an aqueous sample may also be calculated as follows:

Calcium Hardness, mg equivalent CaCO<sub>3</sub>/L = 2.497 (Ca, mg/L)

## KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-615 Revision History Cover Page Page 1

TITLE: DIGESTIC	ON AND ANALYSIS OF AQUEOUS SAMPL 0 7470	ES FOR MERCURY BY USEPA
Prepared By:	George Brewer	Date: OVO I
Approved By:		
Group Supervisor:	Storge Brewer	Date: 01/29/01
Operations Manager	: Jol C. Benton	Date: 1/29/01
QA Officer:	Dutorah J. Kadeau	Date:
General Manager:	Dernau J. hugas	Date: 1   07   07
	, O	<b>8</b>
Revision History:		

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
7470A	NA	Dn	1-29-01	1/09/01
01	Revised Sect. 4, 5 and 7 to reflect current prac- tice. Revised Sect. 8 to reflect current QC limits. levised sect. 10 to reflect current Applicable Documents and references. Removed figure 2. Update table 1 to reflect current QC limits. Minorchanges through out	CA D	02-16-05	02-16-05
ાર	updated Fig. 1 - new preplogbook page	LAT	04/08	04/08
03	Updated Figure 1 - Example of a Mercury Preparation Loghock page	UAN	03109	03/09
04	Added LOD definition. Updated sections 8,9,10 and Table 1 for DOD QSM version 4.1 compliance.	Dr	08/09	08/09

SOP Number: CA-615 Revision History Cover Page Page 1

TITLE: DIGESTION AND ANALYSIS OF AQUEOUS SAMPLES FOR MERCURY BY USEPA METHOD 7470

Revision History:

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
05	Added Table 2 - DoD QSM Version 4.1 QC Requirements.	LAVO	04/10	04/10
04	Sect. 4.4 - Changed thermometer type. Sect. 7.3 - Changed type of morker sed. Table 1 - Add POL Standard corrective action. Table 2 - added Comments for cali bration blank. Sect. 9 - Added MDL, LOD and LOG information	LAD	oslu	0511
07	Sect. 7- Calibration preparent algesting all to discussing high STD. and ailuting down Added Serial divition and PDS to Sect. 8. Added more MDL, LOD & LOQ information to Sect. 9. Updated and added references to Sect. 10	LAY	04/12	04/12

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TITLE:	DIGESTION AND ANALYSIS OF AQUEOUS SAMPLES FOR MERCURY BY USEPA METHOD 7470	
	nowledge receipt of this standard operating procedure by signing and dating both of t vided. Return the bottom half of this sheet to the QA Department.	
	ge receipt of copy of document SOP CA-615-07, titled DIGESTION AND OF AQUEOUS SAMPLES FOR MERCURY BY USEPA METHOD 7470.	
Recipient:	Date:	
	ANALYTICAL SERVICES, INC. O OPERATING PROCEDURE	
	ge receipt of copy of document SOP CA-615-07, titled DIGESTION AND OF AQUEOUS SAMPLES FOR MERCURY BY USEPA METHOD 7470.	
Recipient:	Date:	

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## TITLE: DIGESTION AND ANALYSIS OF AQUEOUS SAMPLES FOR MERCURY BY USEPA METHOD 7470

#### 1.0 SCOPE AND APPLICATION

The purpose of this SOP is to describe the procedure used by Katahdin Analytical Services, Inc. personnel for the digestion and analysis aqueous samples for mercury using cold vapor atomic absorption spectrophotometry.

This method is applicable to the determination of mercury in groundwaters, aqueous wastes, and mobility-procedure extracts under USEPA Method 7470 (<u>Test Methods for Evaluating Solid Wastes: Physical/Chemical Methods</u>, SW-846, 2nd edition, 1982 (revised 1984), 3rd edition, 1986, and Updates I, II, IIA, and III 1996, Office of Solid Waste and Emergency Response, U.S. EPA.

#### 1.1 Definitions

- <u>CCB</u> Continuing Calibration Blank An analyte-free solution consisting of acidified laboratory grade reagent water used to verify calibration accuracy periodically during analysis.
- <u>CCV</u> Continuing Calibration Verification A midrange standard used to verify calibration accuracy periodically during analysis.
- <u>ICB</u> Initial Calibration Blank An analyte-free solution consisting of acidified laboratory grade reagent water used to verify calibration accuracy.
- <u>ICV</u> Initial Calibration Verification A standard made from a source independent from the calibration standards and with analyte concentrations different from those in the CCV; used to verify the accuracy of the instrument calibration.
- <u>PB</u> Preparation Blank Laboratory grade reagent water that has been brought through the sample preparation process.
- <u>LCS</u> Laboratory Control Sample A standard or solid reference material that has been brought through the sample preparation process.

<u>Matrix Spike</u> - An aliquot of a sample to which a known amount of analyte has been added before digestion.

<u>Duplicate</u> - A second aliquot of a sample that is prepared and analyzed in the same way as the original sample in order to determine the precision of the method.

<u>Serial Dilution</u> - The dilution of a sample by a factor of five. When corrected by the dilution factor, the measured analyte concentrations of the diluted sample should agree with those of the original undiluted sample within specified limits. Serial dilution may reflect the influence of interferents.

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# TITLE: DIGESTION AND ANALYSIS OF AQUEOUS SAMPLES FOR MERCURY BY USEPA METHOD 7470

<u>IDL</u> - Instrument Detection Limit - The lowest concentration of an analyte that can be determined with 95% confidence by the instrument.

<u>MDL</u> - Method Detection Limit - The minimum concentration of an analyte that can be identified, measured, and reported with 99% confidence that the analyte concentration is greater than zero.

<u>LOD</u> – Limit of Detection – An estimate of the minimum amount of a substance that an analytical process can reliably detect. An LOD is analyte and matrix-specific and is used for DoD QSM acceptance criteria.

<u>PQL</u> - Practical Quantitation Limit - The lowest concentration of an analyte that is routinely reported by the laboratory; nominally three to five times the IDL.

#### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of analysts experienced in the analysis of mercury by USEPA Method 7470. Each analyst must demonstrate and document their ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, current revision, "Personnel Training & Documentation of Capability".

It is the responsibility of all Katahdin technical personnel involved in analysis of mercury by USEPA Method 7470 to read and understand this SOP, to adhere to the procedures outlined, and to properly document their data in the appropriate lab notebook. Any deviations from the test or irregularities with the samples should also be recorded in the lab notebook and reported to the Department Manager or designated qualified data reviewer responsible for this data.

It is the responsibility of the Department Manager to ensure that members of their group follow this SOP, that their work is properly documented, and to indicate periodic review of the associated logbooks.

#### 1.3 Safety

Many of the samples and reagents used in cold vapor atomic absorption are toxic or corrosive. Rubber gloves, safety glasses, lab coats, and other protective clothing should be worn whenever these materials are handled. Because of the toxic nature of mercury vapor, care must be taken to avoid its inhalation. The instrument exhaust fan must be in operation whenever the mercury analyzer is in use (the fan should never be shut off).

Users of this procedure must be cognizant of inherent laboratory hazards, proper disposal procedures for contaminated materials and appropriate segregation of hazardous wastes. The toxicity or carcinogenicity of each reagent used in this

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# TITLE: DIGESTION AND ANALYSIS OF AQUEOUS SAMPLES FOR MERCURY BY USEPA METHOD 7470

method has not been precisely defined; however, each chemical should be treated as a potential health hazard. A reference file of material safety data sheets is available to all personnel involved in the chemical analysis. Everyone involved with the procedure must be familiar with the MSDSs for all the materials used in this procedure.

Each qualified analyst or technician must be familiar with the Katahdin Analytical Environmental Health and Safety Manual including the Katahdin Hazardous Waste Management Plan and follow appropriate procedures such as wearing safety glasses and gloves when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location and use of all safety equipment.

#### 1.4 Pollution Prevention/Waste Disposal

Whenever possible, laboratory personnel should use pollution prevention techniques to address their waste generation. Refer to the current revision of the Katahdin Hazardous Waste Management Plan for further details on pollution prevention techniques.

Samples, sample digestates, standards, and other reagents used in cold vapor atomic absorption may contain high concentrations of acids, mercury, and other toxic metals. They should be disposed of in a manner appropriate to the types of hazards they present. All digested mercury samples and standards and excess reagents and standards should be disposed of in the satellite waste container for corrosive wastes (labeled "Waste Stream A") that is located in the Metals Prep lab. Further information regarding waste classification and disposal may be obtained by consulting the laboratory's Hazardous Waste Management Plan and Safety Manual and the Department Manager.

#### 2.0 SUMMARY OF METHOD

The cold vapor atomic absorption technique is based on the absorption of radiation at 253.7 nm by mercury vapor. It relies on the volatility of elemental mercury at room temperature. During preparation, organic mercurials are oxidized and elemental mercury is ionized to Hg<sup>3+</sup>. During instrumental analysis, mercuric ions are reduced to elemental mercury by the addition of stannous chloride. Elemental mercury is then aerated from solution and passes through a cell positioned in the path of a mercury spectrophotometer, where absorbance (peak height) is measured as a function of mercury concentration and recorded by the associated computer. The mercury vapor is then swept out of the instrument into an exhaust hood, where it is evacuated from the laboratory.

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# TITLE: DIGESTION AND ANALYSIS OF AQUEOUS SAMPLES FOR MERCURY BY USEPA METHOD 7470

#### 3.0 INTERFERENCES

In addition to inorganic forms of mercury, organic mercurials may be present in environmental samples. These organo-mercury compounds will not respond to the cold vapor atomic absorption technique unless they are first broken down and converted to mercuric ions. The presence of undigested organo-mercurials in samples will result in a low bias for analytical results. Certain volatile organic materials will also non-specifically absorb radiation at the 253.7 nm analytical wavelength. The presence of such compounds may result in a high bias for analytical results. For these reasons, complete digestion using potassium permanganate and potassium persulfate is required for all environmental samples. Complete digestion is indicated by the persistence of the purple permanganate color (indicating the presence of excess permanganate) following digestion.

Sea waters, brines, and industrial effluents high in chlorides may require additional permanganate to maintain a persistent purple color following digestion. During the oxidation step, chlorides are converted to free chlorine which will absorb radiation at the 253.7 nm analytical wavelength. Any free chlorine thus generated will be present in the headspace of the digestion vessel following digestion. Because samples are poured into autosampler tubes prior to analysis by the mercury analyzer, any free chlorine present in the headspace of the digestion vessels is not sampled by the instrument and the analysis is free of chlorine interference.

#### 4.0 APPARATUS AND MATERIALS

- 4.1 40 mL VOA vials, for use as digestion vessels.
- 4.2 250 mL Pyrex media bottles with plastic screw caps, for use in digesting calibration standards.
- 4.3 Water bath capable of maintaining a constant temperature of 95° C.
- 4.4 Adjustable volume automatic pipettes 2 to 20 uL, 10 to 100 uL, 100 to 1000 uL. Calibrated Eppendorf Reference pipets and Finn digital pipets are appropriate.
- 4.5 Repipetters (adjustable repeating pipetters with reservoirs) for dispensing concentrated nitric acid, concentrated sulfuric acid, and other reagents
- 4.6 Battery powered Traceable Pocket-Size Thermometer from Fisher Scientific, NIST-traceable, covering the range from -50° to 750° C, for monitoring the temperature of the water bath. Mercury-filled thermometers are not acceptable for use in the metals laboratory, due to the possibility of breakage and consequent contamination.
- 4.7 Disposable graduated polystyrene sample cups, 200 mL capacity

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- 4.8 CETAC M-6100 automated mercury analyzer and associated peripherals and parts
- 4.9 Disposable graduated dose cups, 30 mL capacity

Refer to Katahdin SOP CA-629, current revision, "Operation and Maintenance of the CETAC M-6100 Automated Mercury Analyzer" for additional required materials.

#### 5.0 REAGENTS

- 5.1 Laboratory grade reagent water mercury-free water meeting the specifications of ASTM Type II water
- 5.2 Concentrated sulfuric acid, trace metals grade
- 5.3 Concentrated nitric acid, trace metals grade
- 5.4 Concentrated hydrochloric acid, trace metal grade
- 5.5 Potassium permanganate solution, 5% w/v: Dissolve 50 g of potassium permanganate in 1 L laboratory grade reagent water. The source reagent should be labeled as suitable for use in mercury determination.
- 5.6 Potassium persulfate solution, 5% w/v: Dissolve 50g of potassium permanganate in 1L laboratory grade reagent water. The source reagent should be labeled as suitable for use in mercury determination.
- 5.7 Sodium chloride hydroxylamine hydrochloride solution: Dissolve 120 g sodium chloride and 120 g hydroxylamine hydrochloride in laboratory grade reagent water and dilute to a final volume of 1 L.
- 5.8 Stannous chloride solution: Add 70 mL concentrated hydrochloric acid to 500 mL of laboratory grade reagent water. Add 100 g stannous chloride and bring to a final volume of 1 L. Mix to dissolve. Reagent should be labeled as suitable for use in mercury determination.
- Intermediate Mercury Standard A: Appropriately dilute a mercury stock standard to obtain a solution containing 1000 ug of mercury per liter in 2% nitric acid. This intermediate standard is used to prepare calibration standards, matrix spikes, CCVs, and laboratory control samples (refer to Section 8). The identity of the stock standard currently used to prepare this intermediate may be obtained by consulting the Standards Preparation Logbook maintained in the Section. Intermediate Mercury Standard A must be prepared fresh monthly ,and disposed of appropriately after use. (Note: the concentrations of all stock standards must be certified by the vendors as traceable to NIST reference materials).

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5.10 Intermediate Mercury Standard B: Appropriately dilute a mercury stock standard to obtain a solution containing 1000 ug of mercury per liter in 2% nitric acid. The source of the stock standard used to prepare Intermediate Mercury Standard B must be distinct from that used to prepare Intermediate Mercury Standard A (i.e. obtained from a separate vendor). Intermediate Mercury Standard B is used to prepare the ICV (refer to Section 8). The identity of the stock standard currently used to prepare this intermediate standard may be obtained by consulting the Standards Preparation Logbook maintained in the Section. Intermediate Mercury Standard B must be prepared fresh monthly, and disposed of appropriately after use.

#### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

Aqueous samples to be analyzed for mercury should be collected and preserved as described in the following table.

Matrix	Container <sup>1</sup>	Collection Volume/ Weight	Preservation/ Treatment	Holding Time
Aqueous (total)	P, G	250 mL	HNO <sub>3</sub> to pH < 2	28 days
Aqueous (dissolved)	P, G	250 mL	HNO <sub>3</sub> to pH < 2	28 days

<sup>&</sup>lt;sup>1</sup> P = polyethylene or G = glass

#### 7.0 PROCEDURES

#### **BOTTLE PREPARATION**

7.1 Mercury digestions are performed in two different types of vessels. Calibration standards, the Initial Calibration Verification (ICV) standard, and the Initial/Continuing Calibration Blank (ICB/CCB) are prepared in 250 mL Pyrex media bottles. Large bottles are used to provide sufficient volumes of these standards to allow for multiple reanalyses when required. Field samples, Method Blanks, and Laboratory Control Samples are digested in 40 mL VOA vials. These smaller vials provide enough digestate to allow one or two reanalyses when required, but reduce the amounts of samples consumed and waste generated.

VOA vials are reused if the samples they have contained have no measurable mercury above the PQL. After the previous contents of the vials have been discarded, these vials are segregated according to whether the measured mercury concentrations of the previous contents were above the PQL (contaminated vials) or below the PQL (uncontaminated vials). Labels are removed from the vials by wiping

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with a paper towel saturated with toluene. Uncontaminated vials are rinsed with laboratory grade reagent water. Contaminated vials are discarded.

The Pyrex media bottles in which standards are prepared are emptied, rinsed, and reused. Each of these bottles is permanently marked with the concentration of the standard it contains.

#### PREPARATION OF STANDARDS, QC SAMPLES, AND BLANKS

- 7.2 Prior to performing the digestion, make a list of the samples that are to be digested. Enter digestion information (Katahdin Sample Numbers, QC Batch ID, preparation date, analyst initials, etc.) into the ACCESS Metals database and print out a copy of the sample prep bench sheet. All necessary details of sample preparation (standards preparation information, digestion times, initial and final volumes, pertinent observations, etc.) must be recorded on this spreadsheet, which will be bound in the Mercury Preparation Logbook. Refer to Figure 1 for an example page from the Mercury Preparation Logbook.
- 7.3 Using an industrial marker with super permanent ink, label clean VOA vials with the appropriate sample numbers and standard identifications for each sample and standard to be digested.
- 7.4 Use a bottle-top dispenser to add 100 mL of laboratory grade reagent water to a standard digestion bottle (250 mL media bottles). Using a calibrated adjustable pipette, prepare the high calibration standard by adding 1000 uL of Intermediate Mercury Standard A to an appropriately labeled media bottle containing 100 mL of laboratory grade reagent water. The mercury concentration of this calibration standard is 10.0 ug/L. Calibration levels 0.2 ug/L, 0.5 ug/L, 1.0 ug/L, 5.0 ug/L are made by diluting the digested 10.0 ug/L standard into calibration blank solution. See below for amounts. The 0.2 ug/L and 5 ug/L standards are analyzed after calibration as the PQL standard and the CCV (refer to Section 8.0), respectively, as well as being used in the creation of the calibration curve.

Calibration Level	Amount added	Amount calibration blank
		solution
0.2 ug/L	0.3 mL	14.7 mL
0.5 ug/L	L 0.5 mL 9.5 mL	
1.0 ug/L	1 mL	9 mL
5.0 ug/L	5 mL	5 mL

7.5 Add 100 mL of laboratory grade reagent water to the media bottle labeled "ICV". Using a calibrated adjustable pipette, prepare the Initial Calibration Verification standard (refer to Section 8) by adding 600 uL of Intermediate Mercury Standard B to the water in this bottle, and record the bottle number in the Mercury Preparation Logbook. The mercury concentration of the ICV is 6.0 ug/L.

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- 7.6 Prepare an appropriate number of preparation blanks (PBW) by adding 25 mL of laboratory grade reagent water to labeled vials.
- 7.7 Prepare an appropriate number of laboratory control samples (LCSW) by adding 125 uL of Intermediate Mercury Standard A to labeled digestion vials containing 25 mL of laboratory grade reagent water. The mercury concentration of each LCSW is 5.0 ug/L.
- 7.8 Matrix spikes are prepared by adding 25 uL of Intermediate Mercury Std A to 25 mL aliquots of samples. The concentration of mercury added to each matrix spike is 1.0 ug/L.
- 7.9 All QC samples and blanks are digested in the same manner as client samples. Refer to Sample Preparation and Digestion, sections 7.10 through 7.13 of this SOP. The volumes of reagents added to the standards prepared in the media bottles are four times those listed in sections 7.10 through 7.13.

#### SAMPLE PREPARATION AND DIGESTION

- 7.10 Using a graduated disposable dosecup, transfer 25 mL of sample, or an aliquot diluted to 25 mL, to a digestion vial. Add 1.25 mL of concentrated sulfuric acid and 0.625 mL of concentrated nitric acid, swirling to mix after each addition. Add 3.75 mL of potassium permanganate solution, swirl to mix, and allow to stand for at least 15 minutes. Samples that contain large amounts of organic substances may require additional 3.75 mL aliquots of potassium permanganate solution. This is indicated by the failure of the purple permanganate color to persist for the entire 15 minute waiting period. Add additional 3.75 mL aliquots to samples as necessary until the purple color persists for 15 minutes. If any of the samples require these additional aliquots of potassium permanganate solution, record the additional volume used for each sample on the mercury preparation benchsheet.
- 7.11 Add 2 mL of potassium persulfate solution to each sample. Cap the vials and place them in a preheated water bath. Monitor the temperature of the bath with a spirit thermometer throughout the digestion. The temperature of the water bath will fall below 95° C upon addition of the digestion vials. After the temperature of the bath has risen back to 95° C, continue heating the samples at 95° C for two hours. Record initial and final digestion times and temperatures in the mercury prepareation benchsheet.
- 7.12 Remove bottles from the water bath and allow to cool to room temperature. If the purple permanganate color has failed to persist after digestion in any of the samples, add additional 3.75 mL aliquots of potassium permanganate solution as required to the samples, and record these additions in the mercury preparation benchsheet. Heat the samples that required additional permanganate in the water bath at 95° C for an additional two hours. Remove the bottles from the water bath

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and allow to cool to room temperature. If the purple color fails to persist after the second heating step, consult the Department Manager for advice on how to proceed.

7.13 Add 1.5 mL of sodium chloride – hydroxylamine hydrochloride solution to each digestion vial and swirl to mix. This will reduce the excess permanganate, and the sample will change from purple to colorless. Wait at least 30 seconds before proceeding with analysis.

#### **INSTRUMENTAL ANALYSIS**

7.14 Digested mercury samples are analyzed using the CETAC M-6100 Automated Mercury Analyzer. Analysis is automated and is controlled by the QuickTrace Mercury Analyzer software running on a dedicated PC. Detailed instructions for setting up the instrument and analyzing samples are given Katahdin SOP CA-629, "Operation and Maintenance of the CETAC M-6100 Automated Mercury Analyzer".

#### METHOD OF STANDARD ADDITIONS

- 7.15 The standard addition technique involves adding known amounts of standard to one or more aliquots of the processed sample solution. This technique compensates for a sample constituent that enhances or depresses the analyte signal, thus producing a different slope from that of the calibration standards. It will not correct for additive interferences which cause a baseline shift. The method of standard additions shall be used for analysis of all EP extracts, on all analyses submitted as part of a delisting petition, and whenever a new sample matrix is being analyzed.
  - 7.15.1 The simplest version of this technique is the single-addition method, in which two identical aliquots of the sample solution, each of volume  $V_x$ , are taken. To the first (labeled A) is added a known volume  $V_S$  of a standard analyte solution of concentration  $C_S$ . To the second aliquot (labeled B) is added the same volume  $V_S$  of the solvent. The analytical signals of A and B are measured and corrected for nonanalyte signals. The unknown sample concentration  $C_x$  is calculated:

$$C_X = \frac{S_B V_S C_S}{(S_A - S_B) V_X}$$

where  $S_A$  and  $S_B$  are the analytical signals (corrected for the blank) of solutions A and B, respectively.  $V_s$  and  $C_s$  should be chosen so that  $S_A$  is roughly twice  $S_B$  on the average, avoiding excess dilution of the sample. If a separation or concentration step is used, the additions are best made first and carried through the entire procedure.

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- 7.15.2 Improved results can be obtained by employing a series of standard additions. To equal volumes of the sample are added a series of standard solutions containing different known quantities of the analyte, and all solutions are diluted to the same final volume. For example, addition 1 should be prepared so that the resulting concentration is approximately 50 percent of the expected absorbance from the endogenous analyte in the sample. Additions 2 and 3 should be prepared so that the concentrations are approximately 100 and 150 percent of the expected endogenous sample absorbance. The absorbance of each solution is determined and then plotted on the vertical axis of a graph, with the concentrations of the known standards plotted on the horizontal axis. When the resulting line is extrapolated to zero absorbance, the point of interception of the abscissa is the endogenous concentration of the analyte in the sample. The abscissa on the left of the ordinate is scaled the same as on the right side, but in the opposite direction from the ordinate. An example of a plot so obtained is shown in Figure 3. A linear regression program may be used to obtain the intercept concentration.
- 7.15.3 For the results of this MSA technique to be valid, the following limitations must be taken into consideration:
  - The apparent concentrations from the calibration curve must be linear over the concentration range of concern. For the best results, the slope of the MSA plot should be nearly the same as the slope of the standard curve. If the slope is significantly different (greater than 20%), caution should be exercised.
  - The effect of the interference should not vary as the ratio of analyte concentration to sample matrix changes, and the standard addition should respond in a similar manner as the analyte.
  - The determination must be free of spectral interference and corrected for nonspecific background interference.

#### DATA REDUCTION AND REPORTING

7.16 Results are obtained in concentration units (ug/L) from the instrument. Electronic instrument data files are imported into the Metals ACCESS database for data reduction. Sample preparation information (initial sample volumes and final digestate volumes) are entered directly into the Metals ACCESS database to allow calculation of final results for reporting. Results are calculated as follows:

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where: MC = Measured mercury concentration (ug/L)

DF = Dilution factor at instrument IV = Initial sample volume (mL) FV = Final digestate volume (mL)

- 7.17 Results that exceed the calibration range of the instrument may not be reported the sample must be appropriately diluted and reanalyzed. Results for diluted samples should be multiplied by the dilution factor prior to reporting. If additional aliquots of potassium permanganate were added during digestion, the resulting dilution must be corrected for before reporting.
- 7.18 Results are reported down to the laboratory's practical quantitation level (PQL), unless otherwise requested. Results below the PQL should be reported to the PQL and flagged with a "U" qualifier.

#### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

USEPA Method 7470 requires the laboratory to perform specific quality control checks to assess laboratory performance and data quality. Minimum frequencies, acceptance criteria, and corrective actions for these control checks are tabulated in Table 1 and are described below. Preparation instructions and the resulting mercury concentrations for calibration standards, QC standards, and matrix spikes are detailed in Sections 7.4 through 7.8 of this SOP. Table 1 criteria are intended to be guidelines for analysts. The table does not cover all possible situations. If any of the QC requirements are outside the recovery ranges listed in Table 1, all associated samples must be evaluated against all the QC. In some cases data may be reported, but may be reanalyzed in other cases. Making new reagents and standards may be necessary if the standardization is suspect. The corrective actions listed in Table 1 may rely on analyst experience to make sound scientific judgments. decisions are based on holding time considerations and client and project specific Data Quality Objectives. The Department Manager, Operations Manager, General Manager and/or Quality Assurance Officer may be consulted to evaluate data. Some samples may not be able to be reanalyzed within hold time. In these cases "qualified" data with narration may be advisable after consultation with the client.

In some cases the standard QC requirements listed in Table 1 may not be sufficient to meet the Data Quality Objectives of the specific project. Much of the work performed at the lab is analyzed in accordance with specific QC requirements spelled out in a project specific Quality Assurance Project Plan (QAPP) or in a program specific Quality Systems Manual (QSM). The reporting limits, acceptance criteria and/or corrective actions may be different than those specified in this SOP. In these cases the appropriate information will be communicated to the Department Manager and/or senior chemists before initiation of the analyses so that specific product codes can be produced for the project. In addition, the work order notes for each project will describe the specific QAPP or QSM to be followed.

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- 8.1 Instrument detection limits (IDL) are determined quarterly for each analyte analyzed on each instrument by each method. This determination requires seven replicate analyses of a laboratory grade reagent water spiked at 3-5 times the anticipated detection limit for each analyte, performed on three non-consecutive days. The standard deviation of the 21 analyses is multiplied by three to obtain the IDL. For more information on performing IDL determinations, refer to the current revision of Katahdin SOP QA-806.
- 8.2 Method detection limits (MDL) are determined annually for each analyte analyzed on each instrument. This determination requires at least seven replicate digestions and analyses of laboratory grade reagent water spiked at 3-5 times the anticipated MDL for each analyte. MDLs differ from IDLs in that the replicates are digested prior to analysis, and they may be analyzed on a single day. The standard deviation of the 7 (or more) replicate analyses is multiplied by the Student's t-value to obtain the MDL. For more information on performing MDL determinations, refer to the current revision of Katahdin SOP QA-806.
- 8.3 Limits of Detection (LOD) are used when evaluating data using DoD QSM. The LOD is established by spiking a quality system matrix at 2-3 times the detection limit for a single analyte standard and 1-4 times the detection limit for a multi-analyte standard. The LOD must be verified quarterly. For more information on performing LOD determinations, refer to the current revision of Katahdin SOP QA-806.
- 8.4 Instrument calibration The instrument must be calibrated each time it is set up, and calibration standards must be digested each day that samples are digested. Calibration includes analysis of a calibration blank and five calibration standards with graduated concentrations in the appropriate range. The concentration of one of the calibration standards must be at the Practical Quantitation Level (PQL). The intermediate standards used for preparing the calibration standards are prepared at least once per month in 2% nitric acid. Because mercury may be adsorbed onto the walls of glass and plastic containers, the calibration standards must be prepared fresh daily. The correlation coefficient for the calibration curve must be at least 0.995. If the calibration curve does not pass this test, analysis must be halted, the problem corrected, and the instrument recalibrated.
- 8.5 An Initial Calibration Verification (ICV) solution is analyzed after the initial calibration to check calibration accuracy. The ICV solution is prepared from a standard source different than that of the calibration standard and at a concentration within the working range of the instrument. The result of the ICV must fall within 90% to 110% of the expected value. If the ICV fails, results may not be reported from the run until the problem is corrected and a passing ICV has been analyzed.
- 8.6 The Continuing Calibration Verification (CCV) solution is analyzed after the initial calibration, after every ten samples, and at the end of the analytical run. The CCV solution is prepared using the same standard used for calibration at a concentration

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near the mid-point of the calibration curve. Results of the CCVs must fall within 80% to 120% of the expected value. If a CCV fails, associated sample results may not be reported from the run until the problem is corrected and a passing CCV has been analyzed. Also, all samples analyzed after the last passing CCV must be reanalyzed.

- 8.7 A calibration blank is analyzed after each ICV and CCV. A calibration blank that is analyzed after the ICV is called an Initial Calibration Blank (ICB). A calibration blank that is analyzed after a CCV is called a Continuing Calibration Blank (CCB). The absolute values of results of ICBs and CCBs must be less than the Practical Quantitation Level (PQL) for each element. If samples are being run using DoD QSM criteria, the absolute values of ICBs and CCBs must be less than the Limit of Detection (LOD). If an ICB or a CCB fails, results for the failing elements may not be reported from the run until the problem is corrected and a passing ICB or CCB has been analyzed. Also, all samples analyzed after the last passing CCB must be reanalyzed.
- 8.8 A standard with a mercury concentration that is at the Practical Quantitation Limit (PQL) is analyzed at the beginning of the run to determine calibration accuracy at the reporting limit. Result of the PQL standard should fall within 70% to 130% of the expected values. No corrective action has been established at this time.

#### PREPARATION BATCH QC SAMPLES

- 8.9 Preparation blank (PBW or PBS), consisting of reagent water carried through the same process as associated samples, is prepared with each digestion batch of twenty or fewer samples. The results of preparation blanks must be less than the Practical Quantitation Level (PQL) for each element. For DoD QSM acceptance criteria the results must be less than ½ the PQL except for common contaminants which must be less than the PQL. If a preparation blank fails, results for the failing elements may not be reported from the digestion batch, and all associated samples must be redigested, with the following exception. If the result for a preparation blank is greater than the PQL (greater than ½ PQL for DoD), associated sample results that are less than the PQL (less than ½ PQL for DoD) or greater than or equal to ten times the measured preparation blank concentration may be reported.
- 8.10 A laboratory control sample (LCSW), consisting of spiked reagent carried through the same process as associated samples, is prepared with each digestion batch of twenty or fewer samples. Results for laboratory control samples must fall within 80% to 120% of the expected value, unless laboratory-generated statistical limits are available. If a laboratory control sample fails, results may not be reported from the digestion batch, and all associated samples must be redigested.

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#### SAMPLE MATRIX QC SAMPLES

8.11 Matrix spiked duplicate samples are prepared at a minimum frequency of one per digestion batch. Matrix spike recoveries for these samples are calculated as follows:

Recovery (%) = 
$$\frac{(P-S)}{A} \times 100\%$$

where: P = Spiked sample value

S = Original sample value

A = Spike amount

The recovery for each element in a spiked sample or spiked duplicate sample must fall within 75% to 125% of the actual value if the result for the unspiked sample is less than four times the amount of spike added. If one or both spike recoveries fail, a matrix interference should be suspected and the associated sample result should be flagged on the report of analysis. If DoD QSM acceptance criteria are being used, recoveries must be the same as stated for laboratory control samples.

The relative percent difference between matrix spiked duplicate sample results is calculated as follows:

RPD (%) = 
$$\frac{|D_1 - D_2|}{(|D_1 + D_2|)/2} \times 100$$

where:  $D_1$  = Spike sample result

D<sub>2</sub>= Spike duplicate sample result

A control limit of 20% RPD is applied to matrix spike duplicate analysis. If the matrix spike duplicate analysis fails, the associated sample result should be flagged on the report of analysis.

A serial dilution is analyzed to check for chemical or physical interferences. If the analyte concentration of a sample is sufficiently high (minimally, 50 x IDL or 50 x LOQ if using DoD QSM acceptance criteria), the measured concentration of a serial dilution (1:5 dilution) of the sample should agree within 90% to 110% of the original determination. The percent difference between the original sample and the serial dilution should be calculated as follows:

Difference (%) = 
$$\frac{|L-S|}{S}$$
 \*100%

where: L = Serial dilution result (corrected for dilution)

S = Original sample result

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If the serial dilution analysis fails, a matrix interference should be suspected. The associated sample result should be flagged on the report of analysis or the sample should be reanalyzed at dilution to eliminate the interference.

8.13 Post-digestion Spike (PDS) additions must be performed for DoD QSM samples if the serial dilution is not within acceptance criteria or if the analyte concentrations in all samples are less than 50x the LOD. The spike addition should produce a concentration that is between 10 and 100x the LOQ. The recovery of the PDS must be within 75-125%. If the PDS fails, all samples must be run by method of standard additions or appropriately flagged.

#### 9.0 METHOD PERFORMANCE

The method detection limit (MDL) is defined as the minimum concentration of a substance that can be measured and reported with 99% confidence that the value is above zero. The MDLs shall be determined and then verified one time per type of instrument.

Limits of Detection (LODs) must be determined for all parameters for which the laboratory is accredited under the DoD Environmental Laboratory Accreditation Program. LOD's must be verifified for every preparation and analytical method combination and on every applicable instrument on a guarterly basis.

The LOQs/PQLs shall be set at the lowest point in the calibration curve for all analyses utilizing an initial calibration. LOQ's must be verified quarterly for every preparation and analytical method combination and on every applicable instrument on a quarterly basis for all parameters included in the DoD Scope of Accreditation.

MDLs are filed with the Organic Department Manager and then with the QAO. LOD and LOQ verifications are filed with the QAO

Refer to the current revision of Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, for procedures on determining the MDL.

Refer to the current revision of USEPA Method 7470 for other method performance parameters and requirements.

#### 10.0 APPLICABLE DOCUMENTS/REFERENCES

Test Methods for Evaluating Solid Wastes, United States Environmental Protection Agency, USEPA SW 846, Third Edition, Final Update III (9/94), Method 7470A.

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Department of Defense Quality Systems Manual for Environmental Laboratories (DOD QSM), Current Version.

The National Environmental Laboratory Accreditation Conference (NELAC) Standards, June 2003.

The NELAC Institute, Laboratory Accreditation Standards, Volume 1, Management and Technical Requirements for Laboratories Performing Environmental Analysis, 10/06/2010.

Katahdin SOP CA-101, Equipment Maintenance and Troubleshooting, current revision.

Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications.

QuickTrace M6100 Mercury Analyzer Operator Manual Version 1.0.1, CETAC Technologies.

QuickTrace Mercury Analyzer Software Manual, CETAC Technologies.

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#### TABLE 1

#### QC REQUIREMENTS

Parameter/ Method	QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action
Mercury/ USEPA 7470	Initial Calibration, 5 points plus a calibration blank.	Daily prior to sample analysis.	Correlation coefficient $\geq$ 0.995.	Correct problem and repeat calibration.
	Initial Calibration Verification (ICV), prepared from a second source.	Before beginning a sample run.	Recovery within <u>+</u> 10% of true value.	Correct problem and repeat calibration.
	Initial Calibration Blank (ICB)	Before beginning a sample run.	Less than PQL.	Correct problem and repeat calibration.
	Practical Quantitation Level Standard (PQL)	Before beginning a sample run.	Recovery within <u>+</u> 30% of true value.	Correct problem and repeat calibration.
	Continuing Calibration Verification (CCV)	At beginning or run, after every 10 samples, and at end of the run	Recovery within <u>+</u> 10% of true value	Repeat calibration and reanalyze all samples analyzed since the last successful CCV.
	Continuing Calibration Blank (CCB)	At beginning or run, after every 10 samples, and at end of the run	Less than PQL.	Repeat calibration and reanalyze all samples analyzed since the last successful CCB.
	Preparation Blank (PBW)	One per digestion batch of 20 or fewer samples.	Less than PQL.	<ul> <li>1) Investigate source of contamination.</li> <li>2) Redigest and reanalyze all associated samples if sample concentration ≥ PQL and &lt; 10x the blank concentration.</li> </ul>
	Laboratory Control Sample (LCSW)	One per digestion batch of 20 or fewer samples.	Recovery within ± 20% of true value.	Redigest all affected samples.
	Matrix Spike Sample (S)	One per digestion batch of 20 or fewer samples.	Recovery ±25% of true value, if sample > 4x spike value.	Flag results.
	Matrix Spike Duplicate Sample (P)	One per digestion batch of 20 or fewer samples.	<ol> <li>Recovery ± 25% of true value, if sample &lt; 4x spike added.</li> <li>RPD ≤20% for duplicate spikes.</li> </ol>	Flag results
	Instrument Detection Limit (IDL) Study	Quarterly.	IDL < PQL	Repeat IDL study.     Raise PQL.
	Limit of Detection (LOD) determination	Quarterly.	LOD = 2-3X MDL	Repeat LOD Determination.
<u>i</u>	Method Detection Limit (MDL) Study		A-806, "Method Detection tudies and Verifications".	n Limit, Instrument Detection Limit, current revision.

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# TABLE 2 DOD QSM VERSION REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Demonstrate acceptable analytical capability	Prior to using any test method and at any time there is a significant change in instrument type, personnel, test method, or sample matrix.	QC acceptance criteria published by DoD, if available; otherwise, method-specified criteria.	Recalculate results; locate and fix problem, then rerun demonstration for those analytes that did not meet criteria.	NA.	This is a demonstration of analytical ability to generate acceptable precision and bias per the procedure in Appendix C. No analysis shall be allowed by analyst until successful demonstration of capability is complete.
LOD determination and verification	(Refer to current revision of SOP QA- 806)				
LOQ establishment and verification	(Refer to current revision of SOP QA- 806)				
Initial calibration (ICAL) for mercury - minimum 5 standards and a calibration blank	Daily ICAL prior to sample analysis.	5 points plus a calibration blank, r ≥ 0.995.	Correct problem, then repeat ICAL.	Flagging criteria are not appropriate.	Problem must be corrected. No samples may be run until ICAL has passed.
Second source calibration verification (ICV)	Once after each ICAL, prior to beginning a sample run.	Value of second source for all analyte(s) within ± 10% of true value.	Correct problem and verify second source standard. Rerun ICV. If that fails, correct problem and repeat ICAL.	Flagging criteria are not appropriate.	Problem must be corrected. No samples may be run until calibration has been verified.
Continuing calibration verification (CCV)	After every 10 field samples and at the end of the analysis sequence.	within ± 20% of true value.	Correct problem, rerun calibration verification. If that fails, then repeat ICAL. Reanalyze all samples since the last successful calibration verification.	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply Q-flag to all results for the specific analyte(s) in all samples since the last acceptable calibration verification.	Problem must be corrected. Results may not be reported without a valid CCV. Flagging is only appropriate in cases where the samples cannot be reanalyzed.

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#### TABLE 2

#### DOD QSM VERSION REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Method blank	One per preparatory batch.	No analytes detected > ½ RL (> RL for common lab contaminants) and > 1/10 the amount measured in any sample or 1/10 the regulatory limit (whichever is greater). Blank result must not otherwise affect sample results.	Correct the problem. Report sample results that are <lod or="">10x the blank concentration. Reprepare and reanalyze the method blank and all associated samples with results &gt; LOD and &lt; 10x the contaminated blank result. Contact Client if samples cannot be reprepped within hold time.</lod>	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply B-flag to all results for the specific analyte(s) in all samples in the associated preparatory batch.	Problem must be corrected. Results may not be reported without a valid method blank. Flagging is only appropriate in cases where the samples cannot be reanalyzed.
Calibration blank	Before beginning a sample run, after every 10 samples, and at end of the analysis sequence.	No analytes detected > LOD.	Correct problem. Reprep and reanalyze calibration blank. All samples following the last acceptable calibration blank must be reanalyzed.	Apply B-flag to all results for specific analyte(s) in all samples associated with the blank.	Problem must be corrected. All samples following the last acceptable calibration blank must be reanalyzed.
LCS	One per preparatory batch.	Water: Recovery must be within + 20% of the true value Soil: Recovery must be within vendor supplied limits (varies by lot).	Correct problem, then reprep and reanalyze the LCS and all samples in the associated preparatory batch for failed analytes, if sufficient sample material is available (see full explanation in Appendix G).	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply Q-flag to specific analyte(s) in all samples in the associated preparatory batch.	Problem must be corrected. Results may not be reported without a valid LCS. Flagging is only appropriate in cases where the samples cannot be reanalyzed.
Matrix spike (MS)	One per preparatory batch per matrix (see Box D-7).	Recovery must be within + 20% of the true value.	Examine the project- specific DQOs. If the matrix spike falls outside of DoD criteria, additional quality control tests are required to evaluate matrix effects.	For the specific analyte(s) in the parent sample, apply J-flag if acceptance criteria are not met.	For matrix evaluation only. If MS results are outside the LCS limits, the data shall be evaluated to determine the source of difference and to determine if there is a matrix effect or analytical error.
Matrix spike duplicate (MSD) or sample duplicate	One per preparatory batch per matrix (see Box D-7).	MSD: Recovery must be within + 20% of the true value. MSD or sample duplicate: RPD ≤ 20% (between MS and MSD or sample and sample duplicate).	Examine the project- specific DQOs. Contact the client as to additional measures to be taken.	For the specific analyte(s) in the parent sample, apply J-flag if acceptance criteria are not met.	The data shall be evaluated to determine the source of difference.

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TITLE: DIGESTION AND ANALYSIS OF AQUEOUS SAMPLES FOR MERCURY BY USEPA METHOD 7470

#### TABLE 2

#### DOD QSM QC REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Method of standard additions (MSA)	When matrix interference is confirmed.	NA.	NA.	NA.	Document use of MSA in the case narrative.
Results reported between DL and LOQ	NA.	NA.	NA.	Apply J-flag to all results between DL and LOQ.	

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TITLE: DIGESTION AND ANALYSIS OF AQUEOUS SAMPLES FOR MERCURY BY USEPA METHOD 7470

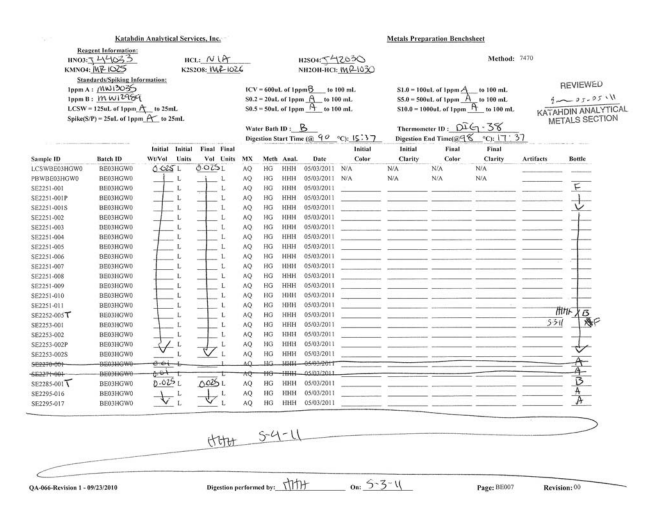
# TABLE 3 SUMMARY OF METHOD MODIFICATIONS

TOPIC	KATAHDIN SOP CA-615-07	USEPA METHOD 7470
Reagents	Stannous chloride dissolved in hydrochloric acid to prevent clogging of mercury analyzer, per instrument manufacturer's recommendation.	Stannous chloride dissolved/suspended in sulfuric acid.
Procedures	1)Sampling and gas stream switching performed automatically by mercury analyzer. 2)Working Mercury standard prepared monthly in 2% nitric; calibration standards prepared fresh daily.	1)Sampling and gas stream switching performed manually by analyst. 2)Working Mercury standard prepared fresh daily and acidity maintained at 0.15% nitric.
QC – Calibration Verification	<ol> <li>Known reference sample (ICV) analyzed daily.</li> <li>Calibration verified after every 10 samples with CCV.</li> </ol>	Known reference sample analyzed quarterly.     Calibration verified after every 20 samples.
QC - Calibration Blanks	Acceptance criteria employed for 245.1: $\pm$ PQL	Acceptance criteria stated in 245.1: ± MDL

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TITLE: DIGESTION AND ANALYSIS OF AQUEOUS SAMPLES FOR MERCURY BY USEPA METHOD 7470

# FIGURE 1 EXAMPLE PAGE FROM MERCURY PREPARATION LOGBOOK

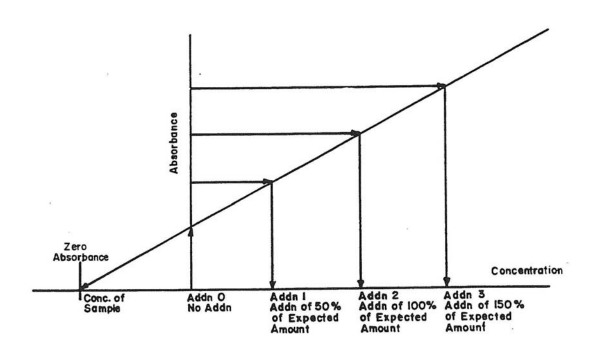


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FIGURE 2 STANDARD ADDITIONS PLOT



# ADDENDUM SOP NO CHANGE FORM

# KATAHDIN ANALYTICAL SERVICES, INC. SOP "REVIEW WITH NO CHANGES" FORM

Name of Person Reviewing SOP: Nicholas 11110	Nost
Review Date: 1/22/13	
SOP Number: CA-615-07	
sop Title: Digestion and analysis of aqu	eous samples for
Mercury by US EPA method 7470	
THE ABOVE REFERENCED SOP HAS BEEN REVIEWED ANALYST OR SUPERVISOR. NO CHANGES ARE REQUI	BY A QUALIFIED AND TRAINED
Department Supervisor Signature:	Date:
- A. Dreyer	02/26/13
QAO Signature:	Date:
Loseio Dirond	030413

# KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-611 Revision History Cover Page Page 1

TITLE:	DIGESTION METHOD 74	AND ANALYSIS OF SOLID SAMPLES FO	R MERC	URY BY USEPA
Prepared E	Зу: _	George Brower	Date:_	12/97
Approved	Ву:			
Group Sup	ervisor: _	Slorge Grewer	Date:_	01/29/01
Operations	s Manager: _	Joh Buta	Date:_	1/29/01
QA Officer	 ·	Deborah J. Nadeau	Date:_	1-29-01
General M	anager: _	Derner F. Kufesh	Date:_	1/29/02
				a de la companya de l
Revision H	listory:			

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
02 7471A	Format changes, added pollution. prevention, other minor changes to sections 7,8 and Qx Table.	On	1.29.01	1/29/01
7471A	Changed Lecman PS200 Automated Mercury Analyzer to Cefac Mc100 Mercury analyzer. Revised Sect. 10 to Show correct reference material. Removed fig. 2 Revised Sect. 4.8, 5.7 and 8.9 to reflect correct practises. Minor changes through out	LAD	031605	021605
04 7471A	Sect. 5:3 and 5:10 - changed preparation of internalid nervoy standards from daily to monthly. Sect. 7.8 - removed each brakish blanks (LCB/CCB). They are prepared in Sect. 7.6. Added weighing of boiling chips for the prep blanks. Sect. 8.3 - Removed intermediate standards	LAD	03/08	80/50
05	Revised Sections 8 and 10, and Tables land 2 to update compliance from method 7471A to method 7471B.	<i>ian</i>	02/09	02/09
06	Added LDD definition. Updated Sections 8, 9,10 and Table 1 for DoD QSM version 4.1 compliance.	Dn	68/09	08/09

# KATAHDIN ANALYTICAL SERVICES, INC. STANDARD OPERATING PROCEDURE

SOP Number: CA-611 Revision History Cover Page (cont.) Page 1

TITLE: DIGESTION AND ANALYSIS OF SOLID SAMPLES FOR MERCURY BY USEPA METHOD 7471

#### Revision History:

SOP Revision	Changes	Approval Initials	Approval Date	Effective Date
٥٦	Added Table 2 with DODOSM Versin 4.1 OC Require ments	LAD	04/10	04/10
08	Sect. 4.6 - Changed thermometer type. Added LCSO-ALCS prepped using agreeds muricing LCSspike. Updated type of marker used to label digestion bottles. Updated corrective action for Guiling PQL Standard.	LAO	12/10	12-110
09	Sect. 7- Changed calibration digestion prom digestion of all points to digestion of high-point and di lution of rest. Changed profifer 320.29 eliquots to 120.69 aliquat Added addition as prop into. Added Serial dilltim and PDS to sect. 8, Added MD, LOQ LOQ into to sect. 9. 4 paded and added references to Sect. 10.	LAVO	oulia	04/12
and the south				

SOP Number: CA-611-09 Date Issued: 04/12 Page 3 of 29

TITLE:	DIGESTION AND ANALYSIS OF SOLID SAMPLES FOR MERCURY BY USEPA METHOD 7471			
Please acknowledge receipt of this standard operating procedure by signing and dating both of the spaces provided. Return the bottom half of this sheet to the QA Department.				
	e receipt of copy of document SOP CA-611-09, Titled Digestion and Analysis of es for Mercury by USEPA Method 7471.			
Recipient:	Date:			
	NALYTICAL SERVICES, INC. OPERATING PROCEDURE			
	e receipt of copy of document SOP CA-611-09, Titled Digestion and Analysis of es for Mercury by USEPA Method 7471.			
Recipient:	Date:			

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# TITLE: DIGESTION AND ANALYSIS OF SOLID SAMPLES FOR MERCURY BY USEPA METHOD 7471

#### 1.0 SCOPE AND APPLICATION

The purpose of this SOP is to describe the procedure used by Katahdin Analytical Services, Inc. personnel for the digestion and analysis solid samples for mercury using cold vapor atomic absorption spectrophotometry.

This method is applicable to the determination of mercury in soils, sediments, bottom deposits, and sludges under USEPA Method 7471 (<u>Test Method for Evaluating Solid Wastes</u>, USEPA SW 846, Third Edition).

#### 1.1 Definitions

- <u>ICB</u> Initial Calibration Blank An analyte-free solution consisting of acidified laboratory reagent grade water used to verify calibration accuracy.
- <u>CCB</u> Continuing Calibration Blank An analyte-free solution consisting of acidified laboratory reagent grade water used to verify calibration accuracy periodically during analysis.
- <u>ICV</u> Initial Calibration Verification A standard made from a source independent from the calibration standards and with analyte concentrations different from those in the CCV; used to verify the accuracy of the instrument calibration.
- <u>CCV</u> Continuing Calibration Verification A midrange standard used to verify calibration accuracy periodically during analysis.
- <u>LCS</u> Laboratory Control Sample A standard or solid reference material that has been brought through the sample preparation process. LCSS utilizes the standard reference material. LCSO is spiked with aqueous mercury LCS spike.
- <u>PB</u> Preparation Blank Laboratory reagent grade water that has been brought through the sample preparation process.

<u>Matrix Spike</u> - An aliquot of a sample to which a known amount of analyte has been added before digestion.

<u>Duplicate</u> - A second aliquot of a sample that is prepared and analyzed in the same way as the original sample in order to determine the precision of the method.

<u>SERIAL DILUTION</u> - The dilution of a sample by a factor of five. When corrected by the dilution factor, the measured analyte concentrations of the diluted sample should agree with those of the original undiluted sample within specified limits. Serial dilution may reflect the influence of interferents.

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### TITLE: DIGESTION AND ANALYSIS OF SOLID SAMPLES FOR MERCURY BY USEPA METHOD 7471

<u>IDL</u> - Instrument Detection Limit - The lowest concentration of an analyte that can be determined with 95% confidence by the instrument.

<u>MDL</u> - Method Detection Limit - The minimum concentration of an analyte that can be identified, measured, and reported with 99% confidence that the analyte concentration is greater than zero.

<u>LOD</u> – Limit of Detection – An estimate of the minimum amount of a substance that an analytical process can reliably detect. An LOD is analyte and matrix-specific and is used for DoD QSM acceptance criteria.

<u>PQL</u> - Practical Quantitation Limit - The lowest concentration of an analyte that is routinely reported by the laboratory; nominally three to five times the IDL.

#### 1.2 Responsibilities

This method is restricted to use by, or under the supervision of analysts experienced in the analysis of mercury by USEPA Method 7471. Each analyst must demonstrate and document their ability to generate acceptable results with this method. Refer to Katahdin SOP QA-805, current revision, "Personnel Training & Documentation of Capability".

It is the responsibility of all Katahdin technical personnel involved in analysis of mercury by USEPA Method 7471 to read and understand this SOP, to adhere to the procedures outlined, and to properly document their data in the appropriate lab notebook. Any deviations from the test or irregularities with the samples should also be recorded in the lab notebook and reported to the Department Manager or designated qualified data reviewer responsible for this data.

It is the responsibility of the Department Manager to ensure that members of their group follow this SOP, to ensure that their work is properly documented, and to initiate periodic review of the associated logbooks.

#### 1.3 Safety

Many of the samples and reagents used in cold vapor atomic absorption are toxic or corrosive. Gloves, safety glasses, lab coats, and other protective clothing should be worn whenever these materials are handled. Because of the toxic nature of mercury vapor, care must be taken to avoid its inhalation. The instrument exhaust fan must be in operation whenever the mercury analyzer is in use (the fan should never be shut off).

Users of this procedure must be cognizant of inherent laboratory hazards, proper disposal procedures for contaminated materials and appropriate segregation of hazardous wastes. The toxicity or carcinogenicity of each reagent used in this

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# TITLE: DIGESTION AND ANALYSIS OF SOLID SAMPLES FOR MERCURY BY USEPA METHOD 7471

method has not been precisely defined; however, each chemical should be treated as a potential health hazard. A reference file of material safety data sheets is available to all personnel involved in the chemical analysis. Everyone involved with the procedure must be familiar with the MSDSs for all the materials used in this procedure.

Each qualified analyst or technician must be familiar with Katahdin Analytical Environmental Health and Safety Manual including the Katahdin Hazardous Waste Plan and must follow appropriate procedures. These include the use of appropriate personal protective equipment (PPE) such as safety glasses, gloves and lab coats when working with chemicals or near an instrument and not taking food or drink into the laboratory. Each analyst should know the location of all safety equipment. Each analyst shall receive a safety orientation from their Department Manager, or designee, appropriate for the job functions they will perform.

#### 1.4 Pollution Prevention/Waste Disposal

Whenever possible, laboratory personnel should use pollution prevention techniques to address there waste generation. Refer to the current revision of the Katahdin Hazardous Waste Management Plan for further details on pollution prevention techniques.

Samples, sample digestates, standards, and other reagents used in cold vapor atomic absorption may contain high concentrations of acids, mercury, and other toxic metals. They should be disposed of in a manner appropriate to the types of hazards they present. All digested mercury samples and standards and excess reagents and standards should be disposed of in the satellite waste container for corrosive wastes (labeled "Waste Stream A") that is located in the Metals Prep lab. Further information regarding waste classification and disposal may be obtained by consulting the laboratory's Katahdin Analytical Environmental Health and Safety Manual and the Department Manager.

#### 2.0 SUMMARY OF METHOD

The cold vapor atomic absorption technique is based on the absorption of radiation at 253.7 nm by mercury vapor. It relies on the volatility of elemental mercury at room temperature. During preparation, organic mercurials are oxidized and elemental mercury is ionized to Hg<sup>3+</sup>. During instrumental analysis, mercuric ions are reduced to elemental mercury by the addition of stannous chloride. Elemental mercury is then aerated from solution and passes through a cell positioned in the path of a mercury spectrophotometer, where absorbance (peak height) is measured as a function of mercury concentration and recorded by the associated computer. The mercury vapor is then swept out of the instrument into an exhaust hood, where it is evacuated from the laboratory.

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#### 3.0 INTERFERENCES

In addition to inorganic forms of mercury, organic mercurials may be present in environmental samples. These organo-mercury compounds will not respond to the cold vapor atomic absorption technique unless they are first broken down and converted to mercuric ions. The presence of undigested organo-mercurials in samples will result in a low bias for analytical results. Certain volatile organic materials will also non-specifically absorb radiation at the 253.7 nm analytical wavelength. The presence of such compounds may result in a high bias for analytical results. For these reasons, complete digestion using potassium permanganate is required for all environmental samples. Complete digestion is indicated by the persistence of the purple permanganate color (indicating the presence of excess permanganate) following digestion.

Samples that are high in chlorides may require additional permanganate to maintain a persistent purple color following digestion. During the oxidation step, chlorides are converted to free chlorine, which will absorb radiation at the 253.7 nm analytical wavelength. Any free chlorine thus generated will be present in the headspace of the digestion vessel following digestion. Because samples are poured into autosampler tubes prior to analysis by the mercury analyzer, any free chlorine present in the headspace of the digestion vessels is not sampled by the instrument and the analysis is free of chlorine interference.

#### 4.0 APPARATUS AND MATERIALS

- 4.1 250 mL Pyrex media bottles with plastic screw caps, for use as digestion vessels.
- 4.2 Water bath capable of maintaining a constant temperature of 95° C.
- 4.3 Analytical balance capable of weighing to 0.01 g.
- 4.4 Adjustable volume automatic pipettes 2 to 20 uL, 10 to 100 uL, 100 to 1000 uL. Calibrated Eppendorf Reference pipets and Finn digital pipets are appropriate.
- 4.5 Repipetters (adjustable repeating pipetters with reservoirs) for dispensing concentrated nitric acid, concentrated sulfuric acid, and other reagents.
- 4.6 Battery powered Traceable Pocket-Size Thermometer from Fisher Scientific, NIST-traceable, covering the range from -50° to 750° C, for monitoring the temperature of the water bath. Mercury-filled thermometers are not acceptable for use in the metals laboratory, due to the possibility of breakage and consequent contamination.
- 4.7 Disposable graduated polystyrene sample cups, 200 mL capacity.

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4.8 CETAC M6100 Mercury Analyzer and associated peripherals and parts.

Refer to Katahdin SOP CA-629, current revision, "Operation and Maintenance of the CETAC M6100 Mercury Analyzer" for additional required materials.

#### 5.0 REAGENTS

- 5.1 Laboratory reagent grade water mercury-free water.
- 5.2 Concentrated nitric acid (HNO<sub>3</sub>), trace metal grade
- 5.3 Concentrated hydrochloric acid (HCI), trace metal grade
- 5.4 Aqua regia: Prepare an appropriate amount immediately before use by carefully adding three volumes of concentrated HCl to one volume of concentrated HNO $_3$  in a heat-proof beaker or flask. Preparation of aqua regia must be performed in a fume hood.
- 5.5 Potassium permanganate solution, 5% w/v: Dissolve 50 g of potassium permanganate in 1 L laboratory reagent grade water. The source reagent should be labeled as suitable for use in mercury determination.
- 5.6 Sodium chloride hydroxylamine hydrochloride solution: Dissolve 120 g sodium chloride and 120 g hydroxylamine hydrochloride in laboratory reagent grade water and dilute to a final volume of 1 L.
- 5.7 Stannous chloride solution: Add 70 mL concentrated hydrochloric acid to 500 mL of laboratory reagent grade water. Add 100 g stannous chloride and bring to a final volume of 1 L. Mix to dissolve. Reagent should be labeled as suitable for use in mercury determination.
- 5.8 Mercury Stock Standards: Two 10.0 mg/L mercury stock standards, obtained from separate sources, are required. The mercury concentrations of these standards must be certified by the manufacturers as traceable to NIST reference standards.
- 5.9 Intermediate Mercury Standard A: Appropriately dilute a mercury stock standard to obtain a solution containing 1000 ug of mercury per liter in 2% nitric acid. This intermediate standard is used to prepare calibration standards, matrix spikes, CCVs, and laboratory control samples (refer to Section 8). The identity of the stock standard currently used to prepare this intermediate standard and instructions for its dilution may be obtained by consulting the Standards Preparation Logbook maintained in the Section. Intermediate Mercury Standard A must be prepared monthly, and disposed of appropriately after use. (Note: the concentrations of all

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# TITLE: DIGESTION AND ANALYSIS OF SOLID SAMPLES FOR MERCURY BY USEPA METHOD 7471

stock standards must be certified by the vendors as traceable to NIST reference materials).

- 5.10 Intermediate Mercury Standard B: Appropriately dilute a mercury stock standard to obtain a solution containing 1000 ug of mercury per liter in 2% nitric acid. The source of the stock standard used to prepare Intermediate Mercury Standard B must be distinct from that used to prepare Intermediate Mercury Standard A (i.e. obtained from a separate vendor). Intermediate Mercury Standard B is used to prepare the ICV (refer to Section 8.0). The identity of the stock standard currently used to prepare this intermediate standard and instructions for its dilution may be obtained by consulting the Standards Preparation Logbook maintained in the Section. Intermediate Mercury Standard B must be prepared monthly, and disposed of appropriately after use.
- 5.11 Solid Reference Material: A soil with a known or empirically-established mercury concentration for use in preparing the laboratory control sample for soils. Solid reference materials should be purchased with certificates listing reference values and quality control acceptance limits. See Figure 3 for an example certificate of analysis for a solid reference material.

#### 6.0 SAMPLE COLLECTION, PRESERVATION AND HANDLING

Soil samples to be analyzed for mercury should be collected and preserved as described in the following table.

Matrix	Container <sup>1</sup>	Collection Volume/ Weight	Preservation/ Treatment	Holding Time
Solid	P, G	40 g	Cool to 4°C ± 2°	28 days

<sup>&</sup>lt;sup>1</sup> P = polyethylene, G = glass

#### 7.0 PROCEDURES

#### **BOTTLE PREPARATION**

7.1 Mercury digestion bottles are reused, and must be cleaned between uses. After the previous contents of the bottles have been discarded, bottles are segregated according to whether the measured mercury concentrations of the previous contents were above the PQL (contaminated bottles) or below the PQL (uncontaminated bottles). Labels are removed from the bottles by wiping with a paper towel saturated with toluene. Both contaminated and uncontaminated bottles are then cleaned with Liquinox and water, if necessary, to remove visible grime, and rinsed thoroughly with tap water.

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- 7.2 Uncontaminated bottles are then triple-rinsed with laboratory reagent grade water, and are ready for reuse.
- 7.3 Contaminated bottles are placed in a bath containing 10% HCl for at least 12 hours. After acid-leaching, these bottles are triple rinsed with laboratory reagent grade water, and are then ready for reuse.

#### PREPARATION OF STANDARDS, QC SAMPLES, AND BLANKS

- 7.4 Prior to performing the digestion, make a list of the samples that are to be digested. Enter digestion information (Katahdin Sample Numbers, Bottle IDs, QC Batch ID, preparation date, analyst initials, etc.) into the ACCESS computer database and print out a copy of the benchsheet. All necessary details of sample preparation (standards preparation information, digestion times, digestion temps, initial weights and final volumes, pertinent observations, etc.) must be recorded on this benchsheet, which will be bound in the Mercury Preparation Logbook. Refer to Figure 1 for an example page from the Mercury Preparation Logbook.
- 7.5 Using an industrial marker with super permanent ink, label clean digestion bottles with the appropriate sample numbers and standard identifications for each sample and standard to be digested.
- 7.6 Use a bottle-top dispenser to add 100 mL of laboratory grade reagent water to a standard digestion bottle (250 mL media bottles). Using a calibrated adjustable pipette, prepare the high calibration standard by adding 1000 uL of Intermediate Mercury Standard A to an appropriately labeled media bottle containing 100 mL of laboratory grade reagent water. The mercury concentration of this calibration standard is 10.0 ug/L. Calibration levels 0.2 ug/L, 0.5 ug/L, 1.0 ug/L, 5.0 ug/L are made by diluting the digested 10.0 ug/L standard into calibration blank solution. See below for amounts. The 0.2 ug/L and 5.0 ug/L standards are analyzed after calibration as the PQL standard and the CCV (refer to Section 8.0), respectively, as well as being used in the creation of the calibration curve.

Calibration Level	Amount added	Amount Calibration Blank Solution
0.2 ug/L	0.3 mL	14.7 mL
0.5 ug/L	0.3 mL	9.5 mL
1.0 ug/L	1 mL	9 mL
5.0 ug/L	5 mL	5 mL

7.7 Using a calibrated adjustable pipette, prepare the initial calibration verification (ICV) standard (refer to Section 8) by adding 600 uL of Intermediate Mercury Standard B to an appropriately labeled digestion bottle. The mercury concentration of the ICV will be 6.0 ug/L.

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- 7.8 Prepare an appropriate number of preparation blanks (PBS) by adding 1.0 g of Teflon boiling chips to labeled digestion bottles.
- 7.9 Prepare an appropriate number of laboratory control samples (LCSS or LCSO) by weighing appropriate masses of solid reference material or by adding 500 uL of Intermediate Mercury Standard A respectively into labeled digestion bottles. The mercury concentration of the LCSS will depend on the solid reference material used, and the mass of each aliquot. Refer to Figure 3 for an example certificate of analysis for a solid reference material. The mercury concentration of the LCSO will be 5.0 ug/L.
- 7.10 Matrix spikes are prepared by adding 100 uL of Intermediate Mercury Std A to each matrix spike sample. The amount of mercury added to each matrix spike increases the final digestate concentration by 1.0 ug/L.
- 7.11 All calibration standards, QC samples, and blanks are digested in the same manner as client samples. Refer to Sample Preparation and Digestion, Steps 7.12 through 7.16 of this SOP.

#### SAMPLE PREPARATION AND DIGESTION

- 7.12 Do not decant any water on the sediment sample. Mix sample with a wooden spatula to ensure homogeneity of the sample. Please refer to the current revision of Katahdin Analytical Services SOP CA-108, "Basic Laboratory Technique", for more detailed guidance on sub-sampling to ensure reproducibility.
  - Weigh an approximate 0.6 g portion of untreated, homogenized sample from the sample container and place in the bottom of a labeled digestion bottle.
- 7.13 Add 5 mL of laboratory reagent grade water and 5 mL of aqua regia to each sample, standard, and QC sample. Place bottles in a water bath located in a fume hood and heat for 2 minutes at 95° C. Remove the bottles from the water bath and allow them to cool in a fume hood.
- 7.14 Add 50 mL of laboratory reagent grade water and 15 mL of potassium permanganate solution to each digestion bottle, swirl to mix, and allow to stand for at least 15 minutes. Samples that contain large amounts of oxidizable organic matter may require additional 15 mL aliquots of potassium permanganate solution. This is indicated by the failure of the purple permanganate color to persist for the entire 15 minute waiting period. Add additional 15 mL aliquots to samples as necessary until the purple color persists for 15 minutes. If any of the samples requires these additional aliquots of permanganate, note that fact on the mercury preparation benchsheet and accordingly adjust the final volumes recorded on the benchsheet for those samples.

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When a persistent purple color has been obtained for all samples, place the digestion bottles in the water bath and heat for 30 minutes at 95° C. Record initial and final time and temperatures on the mercury preparation benchsheet.

- 7.15 Remove the bottles from water bath and allow them to cool in a fume hood. If any of the samples have become colorless during heating, add additional 15 mL aliquots of potassium permanganate solution as necessary to obtain a persistent purple color and heat for an additional 30 minutes at 95° C. Record any information regarding additional permanganate aliquots on the mercury preparation benchsheet and accordingly adjust the final volumes recorded on the benchsheet for the samples affected.
- 7.16 Add 6 mL of sodium chloride hydroxylamine hydrochloride solution to each digestion bottle and swirl to mix. Perform this addition in a fume hood, as chlorine gas may be evolved. This will reduce the excess permanganate, and the sample will change from purple to colorless. Add 50 mL of laboratory reagent grade water to each bottle. Wait at least 30 seconds before proceeding with analysis.

#### **INSTRUMENTAL ANALYSIS**

- 7.17 Digested mercury samples are analyzed using the CETAC M6100 Mercury Analyzer. Analysis is automated and is controlled by the QuickTrace software running on a dedicated PC. Detailed instructions for setting up the instrument and running samples are given Katahdin SOP CA-629, "Operation and Maintenance of the CETAC M6100 Mercury Analyzer". The following information specifically pertains to analysis of digested samples in accordance with USEPA Method 7471, and should be used in conjunction with the instructions given in Katahdin SOP CA-629.
- 7.18 Instrument operating conditions and quality control acceptance limits are specified in the instrument software in "templates". The template that is used to analyze digested samples in accordance with USEPA Method 7471 is named "SW846-7470-7471".
- 7.19 Prior to analysis, digested samples, standards, and QC samples are decanted into autosampler tubes which are placed in racks on the instrument's autosampler. The "standards" autosampler rack has 10 positions for 25 x 100 mm autosampler tubes (50 mL capacity). Tubes containing the calibration standards, the ICV, the CCV, the ICB/CCB, and the PQL standard are placed in the appropriately labeled positions in this autosampler rack.
- 7.20 Client samples, batch QC samples (preparation blanks and laboratory control samples), and matrix QC samples (duplicates and matrix spikes) are decanted into 17 x 100 mm autosampler tubes (15 mL capacity), which are placed in the one of

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the "samples" autosampler racks. The "samples" autosampler racks have 60 positions for 17 x 100 mm autosampler tubes. Instructions for filling the "samples" autosampler racks, including recording the rack position of each sample, are contained in Katahdin SOP CA-629, "Operation and Maintenance of the CETAC M6100 Mercury Analyzer".

#### METHOD OF STANDARD ADDITIONS

- 7.21 The standard addition technique involves adding known amounts of standard to one or more aliquots of the processed sample solution. This technique compensates for a sample constituent that enhances or depresses the analyte signal, thus producing a different slope from that of the calibration standards. It will not correct for additive interferences that cause a baseline shift. The method of standard additions shall be used for analysis of all EP extracts, on all analyses submitted as part of a delisting petition, and whenever a new sample matrix is being analyzed.
  - 7.21.1 The simplest version of this technique is the single-addition method, in which two identical aliquots of the sample solution, each of volume  $V_x$ , are taken. To the first (labeled A) is added a known volume  $V_S$  of a standard analyte solution of concentration  $C_S$ . To the second aliquot (labeled B) is added the same volume  $V_S$  of the solvent. The analytical signals of A and B are measured and corrected for nonanalyte signals. The unknown sample concentration  $C_x$  is calculated:

$$C_X = \frac{S_B V_S C_S}{(S_A - S_B)V_X}$$

where  $S_A$  and  $S_B$  are the analytical signals (corrected for the blank) of solutions A and B, respectively.  $V_s$  and  $C_s$  should be chosen so that  $S_A$  is roughly twice  $S_B$  on the average, avoiding excess dilution of the sample. If a separation or concentration step is used, the additions are best made first and carried through the entire procedure.

7.21.2 Improved results can be obtained by employing a series of standard additions. To equal volumes of the sample are added a series of standard solutions containing different known quantities of the analyte, and all solutions are diluted to the same final volume. For example, addition 1 should be prepared so that the resulting concentration is approximately 50 percent of the expected absorbance from the endogenous analyte in the sample. Additions 2 and 3 should be prepared so that the concentrations are approximately 100 and 150 percent of the expected endogenous sample absorbance. The absorbance of each solution is determined and then plotted on the vertical axis of a graph, with the concentrations of the known standards plotted on the horizontal axis. When the resulting line is

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extrapolated to zero absorbance, the point of interception of the abscissa is the endogenous concentration of the analyte in the sample. The abscissa on the left of the ordinate is scaled the same as on the right side, but in the opposite direction from the ordinate. An example of a plot so obtained is shown in Figure 2. A linear regression program may be used to obtain the intercept concentration.

- 7.21.3 For the results of this MSA technique to be valid, the following limitations must be taken into consideration:
  - The apparent concentrations from the calibration curve must be linear over the concentration range of concern. For the best results, the slope of the MSA plot should be nearly the same as the slope of the standard curve. If the slope is significantly different (greater than 20%), caution should be exercised.
  - The effect of the interference should not vary as the ratio of analyte concentration to sample matrix changes, and the standard addition should respond in a similar manner as the analyte.
  - The determination must be free of spectral interference and corrected for nonspecific background interference.

#### DATA REDUCTION AND REPORTING

7.22 Results are obtained in units of ug/L in the digestate. Results that exceed the calibration range of the instrument may not be reported - the sample must be appropriately diluted and reanalyzed. Results for diluted samples must be multiplied by the dilution factor prior to reporting. If additional aliquots of potassium permanganate were added during digestion, the change in digestate final volume must be taken into account in calculating the final result. Mercury results for solid samples are reported in units of ug/g, calculated on a dry weight basis. Calculation of mercury results for solid samples is performed automatically by the Metals reporting database, as follows:

Mercury Concentration in Solid (mg/kg dry wt.) =  $\frac{(C) \times (DF) \times (FV) \times 100}{(W) \times (TS)}$ 

where C = Measured digestate concentration (ug/L)

DF = Instrument dilution factor FV = Digestate final volume (L)

W = Digested wet sample weight (g)

TS = Total Solids (%)

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7.23 Results are reported down to the laboratory's practical quantitation level (PQL), unless otherwise requested. Results below the PQL should be reported as "<PQL".

#### 8.0 QUALITY CONTROL AND ACCEPTANCE CRITERIA

USEPA Method 7471 requires the laboratory to perform specific quality control checks to assess laboratory performance and data quality. Minimum frequencies, acceptance criteria, and corrective actions for these control checks are tabulated in Table 1 and are described below. Preparation instructions and the resulting mercury concentrations for calibration standards. QC standards, and matrix spikes are detailed in Sections 7.6 through 7.10 of this SOP. Table 1 criteria are intended to be guidelines for analysts. The table does not cover all possible situations. If any of the QC requirements are outside the recovery ranges listed in Table 1, all associated samples must be evaluated against all the QC. In some cases data may be reported, but may be reanalyzed in other cases. Making new reagents and standards may be necessary if the standardization is suspect. The corrective actions listed in Table 1 may rely on analyst experience to make sound scientific judgments. These decisions are based on holding time considerations and client and project specific Data Quality Objectives. The supervisor, Operations Manager, General Manager and/or Quality Assurance Officer may be consulted to evaluate data. Some samples may not be able to be reanalyzed within hold time. In these cases "qualified" data with narration may be advisable after consultation with the client.

Much of the work performed at the lab is analyzed in accordance with specific QC requirements spelled out in a project specific Quality Assurance Project Plan (QAPP) or in a program specific Quality Systems Manual (QSM). The reporting limits, acceptance criteria and/or corrective actions may be different than those specified in this SOP. In these cases the appropriate information will be communicated to the Department Manager and/or senior chemists before initiation of the analyses so that specific product codes can be produced for the project. In addition, the work order notes for each project will describe the specific QAPP or QSM to be followed.

#### INITIAL DEMONSTRATION OF PERFORMANCE

- 8.1 Instrument detection limits (IDL) are determined quarterly for each analyte analyzed on each instrument by each method. This determination requires seven replicate analyses of laboratory reagent grade water spiked, performed on three non-consecutive days. The standard deviation of the 21 analyses is multiplied by three to obtain the IDL. For more information on performing IDL determinations, refer to the current revision of Katahdin SOP QA-806.
- 8.2 Method detection limits (MDL) are determined annually for each analyte analyzed on each instrument. This determination requires at least seven replicate digestions and analyses of laboratory reagent grade water spiked at 3-5 times the anticipated MDL for each analyte. MDLs differ from IDLs in that the replicates are digested

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prior to analysis, and they may be analyzed on a single day. The standard deviation of the 7 (or more) replicate analyses is multiplied by the Student's t-value to obtain the MDL. For more information on performing MDL determinations, refer to the current revision of Katahdin SOP QA-806.

8.3 Limits of Detection (LOD) are used when evaluating data using DoD QSM. The LOD is established by spiking a quality system matrix at 2-3 times the detection limit for a single analyte standard and 1-4 times the detection limit for a multi-analyte standard. The LOD must be verified quarterly. For more information on performing LOD determinations, refer to the current revision of Katahdin SOP QA-806.

#### ANALYTICAL RUN QC

- 8.4 Instrument calibration The instrument must be calibrated each time it is set up, and calibration standards must be digested each day that samples are digested. Calibration includes analysis of a calibration blank and five calibration standards with graduated concentrations in the appropriate range. The concentration of one of the calibration standards must be at the Practical Quantitation Level (PQL). The correlation coefficient for the calibration curve must be at least 0.995. If the calibration curve does not pass this test, analysis must be halted, the problem corrected, and the instrument recalibrated.
- 8.5 An Initial Calibration Verification (ICV) solution is analyzed after the initial calibration to check calibration accuracy. The ICV solution is prepared from a standard source different than that of the calibration standard and at a concentration within the working range of the instrument. The result of the ICV must fall within 90% to 110% of the expected value. If the ICV fails, results may not be reported from the run until the problem is corrected and a passing ICV has been analyzed.
- 8.6 The Continuing Calibration Verification (CCV) solution is analyzed after the initial calibration, after every ten samples, and at the end of the analytical run. The CCV solution is prepared using the same standard used for calibration at a concentration near the mid-point of the calibration curve. Results of the CCVs must fall within 90% to 110% of the expected value. If a CCV fails, associated sample results may not be reported from the run until the problem is corrected and a passing CCV has been analyzed. Also, all samples analyzed after the last passing CCV must be reanalyzed. For DoD QSM acceptance criteria, samples that are below the reporting limit may be reported if the CCV reads greater than 120%.
- 8.7 A calibration blank is analyzed after each ICV and CCV. A calibration blank that is analyzed after the ICV is called an Initial Calibration Blank (ICB). A calibration blank that is analyzed after a CCV is called a Continuing Calibration Blank (CCB). The absolute values of results of ICBs and CCBs must be less than the Practical Quantitation Level (PQL) for each element. If samples are being run using DoD QSM criteria, the absolute values of ICBs and CCBs must be less than the Limit of

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Detection (LOD). If an ICB or a CCB fails, results for the failing elements may not be reported from the run until the problem is corrected and a passing ICB or CCB has been analyzed. Also, all samples analyzed after the last passing CCB must be reanalyzed.

A standard with a mercury concentration that is at the Practical Quantitation Limit (PQL) is analyzed at the beginning of the run to determine calibration accuracy at the reporting limit. Result of the PQL standard should fall within 70% to 130% of the expected values. If the PQL fails, results may not be reported from the run until the problem is corrected and a passing PQL has been analyzed.

#### PREPARATION BATCH QC SAMPLES

- 8.9 Preparation blank (PBW or PBS), consisting of reagent water carried through the same process as associated samples, is prepared with each digestion batch of twenty or fewer samples. The results of preparation blanks must be less than the Practical Quantitation Level (PQL) for each element. For DoD QSM acceptance criteria the results must be less than ½ the PQL except for common contaminants which must be less than the PQL. If a preparation blank fails, results for the failing elements may not be reported from the digestion batch, and all associated samples must be redigested, with the following exception. If the result for a preparation blank is greater than the PQL (greater than ½ PQL for DoD), associated sample results that are less than the PQL (less than ½ PQL for DoD) or greater than or equal to ten times the measured preparation blank concentration may be reported.
- 8.10 A laboratory control sample (LCSS or LCSO), consisting of solid reference material or 500 uL of Intermediate Standard A carried through the same process as associated samples, is prepared with each digestion batch of twenty or fewer samples. If a laboratory control sample fails, results may not be reported from the digestion batch, and all associated samples must be redigested. The laboratory uses a reference value and statistical acceptance limits for laboratory control samples that are supplied by the vendor of the solid reference material. The results of the LCSO must fall with in 80% 120% of its true value which is 5.0 ug/L. If samples are being prepared using DoD QSM acceptance criteria, the results of the LCSO must be within 80% 120%.

#### SAMPLE MATRIX QC SAMPLES

8.11 Matrix spiked duplicate samples are prepared at a minimum frequency of one per digestion batch. Matrix spike recoveries for these samples are calculated as follows:

Recovery (%) = 
$$\frac{(P-S)}{A}$$
 x100%

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where: P = Spiked sample value

S = Original sample value

A = Spike amount

The recovery for each element in a spiked sample or spiked duplicate sample must fall within 75% to 125% of the actual value if the result for the unspiked sample is less than four times the amount of spike added. If one or both spike recoveries fail, a matrix interference should be suspected and the associated sample result should be flagged on the report of analysis. If DoD QSM acceptance criteria are being used, recoveries must be the same as stated for laboratory control samples.

The relative percent difference between matrix spiked duplicate sample results is calculated as follows:

RPD (%) = 
$$\frac{|D_1 - D_2|}{(|D_1 + D_2|)/2} \times 100$$

where:  $D_1$  = Spike sample result

D<sub>2</sub>= Spike duplicate sample result

A control limit of 20% RPD is applied to matrix spike duplicate analysis. If the matrix spike duplicate analysis fails, the associated sample result should be flagged on the report of analysis.

8.12 Serial Dilution - A serial dilution is analyzed to check for chemical or physical interferences. If the analyte concentration of a sample is sufficiently high (minimally, 50 x IDL or 50 x LOQ if using DoD QSM acceptance criteria), the measured concentration of a serial dilution (1:5 dilution) of the sample should agree within 90% to 110% of the original determination. The percent difference between the original sample and the serial dilution should be calculated as follows:

Difference (%) = 
$$\frac{|L-S|}{S}$$
 \*100%

where: L = Serial dilution result (corrected for dilution)

S = Original sample result

If the serial dilution analysis fails, a matrix interference should be suspected. The associated sample result should be flagged on the report of analysis or the sample should be reanalyzed at dilution to eliminate the interference.

8.13 Post-digestion Spike (PDS) additions must be performed for DoD QSM samples if the serial dilution is not within acceptance criteria or if the analyte concentrations in all samples are less than 50x the LOD. The spike addition should produce a concentration that is between 10 and 100x the LOQ. The recovery of the PDS must

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be within 75-125%. If the PDS fails, all samples must be run by method of standard additions or appropriately flagged.

#### 9.0 METHOD PERFORMANCE

The method detection limit (MDL) is defined as the minimum concentration of a substance that can be measured and reported with 99% confidence that the value is above zero. The MDLs shall be determined and verified one time per type of instrument unless otherwise required by the method.

A Limit of Detection (LOD) is an estimate of the minimum amount of a substance that an analytical process can reliably detect. An LOD is analyte and matrix specific and may be laboratory-dependent. LODs must be determined for all parameters for which the laboratory is accredited under the DoD Environmental Laboratory Accreditation Program. LOD's must be verified for every preparation and analytical method combination and on every applicable instrument on a quarterly basis.

The Limit of Detection (LOQ) is the minimum levels, concentrations, or quantities of a target variable (e.g., target analyte) that can be reported with a specified degree of confidence. The LOQ shall be set at the lowest point in the calibration curve for all analyses utilizing an initial calibration. LOQ's must be verified quarterly for every preparation and analytical method combination and on every applicable instrument on a quarterly basis for all parameters included in the DoD Scope of Accreditation. The LOQ must be verified at least once annually if the analysis is not included in the DoD Scope of Accreditation.

MDLs are filed with the Organic Department Manager and then with the QAO. LOD and LOQ verifications are filed with the QAO

Refer to the current revision of Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, for procedures on determining the MDL.

Refer to the current revision of Method 7471 for other method performance parameters and requirements.

#### 10.0 APPLICABLE DOCUMENTS/REFERENCES

Test Methods for Evaluating Solid Waste, Physical/Chemical Methods, USEPA SW846, 3<sup>rd</sup> Edition, Final Updates I, II, IIA, IIB, III, IIIA, IIB and IV, February 2007, Method 7471B.

Department of Defense Quality Systems Manual for Environmental Laboratories (DOD QSM), Current Version.

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The National Environmental Laboratory Accreditation Conference (NELAC) Standards, June 2003.

The NELAC Institute, Laboratory Accreditation Standards, Volume 1, Management and Technical Requirements for Laboratories Performing Environmental Analysis, 10/06/2010.

Katahdin SOP CA-101, Equipment Maintenance and Troubleshooting, current revision.

Katahdin SOP QA-806, Method Detection Limit, Instrument Detection Limit and Reporting Limit Studies and Verifications, current revision.

QuickTrace M6100 Mercury Analyzer Operator Manual Version 1.0.1, CETAC Technologies

QuickTrace Mercury Analyzer Software Manual, CETAC Technologies.

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#### TABLE 1

#### QC REQUIREMENTS

Parameter/ Method		Minimum Frequency	Acceptance Criteria	Corrective Action
Mercury/ USEPA Method 7471B	Initial Calibration, 5 points plus a calibration blank.	Daily prior to sample analysis.	Correlation coefficient ≥ 0.995.	Correct problem and repeat calibration.
		Before beginning a sample run.	Recovery within <u>+</u> 10% of true value.	Correct problem and repeat calibration.
	Initial Calibration Blank (ICB)	Before beginning a sample run.		Correct problem and repeat calibration.
	Practical Quantitation Level Standard (PQL)		Recovery within <u>+</u> 30% of true value.	Correct problem and repeat calibration.
	` ,	At beginning or run, after every 10 samples, and at end of the run	value	Repeat calibration and reanalyze all samples analyzed since the last successful CCV.
	(CCB)	At beginning or run, after every 10 samples, and at end of the run		Repeat calibration and reanalyze all samples analyzed since the last successful CCB.
	Preparation Blank (PBS)	One per digestion batch of 20 or fewer samples.	Less than PQL.	<ol> <li>Investigate source of contamination.</li> <li>Redigest and reanalyze all associated samples if sample concentration ≥ PQL and &lt; 10x the blank concentration.</li> </ol>
	Laboratory Control Sample (LCSS or LCSO)	One per digestion batch of 20 or fewer samples.	LCSS: Recovery within vendor- supplied acceptance limits. LCSO: Recovery within <u>+</u> 20% of true value.	Redigest all affected samples.
	Matrix Spike Sample (S)	batch of 20 or fewer samples.	Recovery ± 25% of true value, if sample > 4x spike value.	Flag results.
	(P) or sample duplicate (D)	batch of 20 or fewer samples.	1)Recovery <u>+</u> 25% of true value, if sample < 4x spike added. 2) RPD ≤20% for duplicate spikes or duplicate samples.	Flag results
	Post-Digestion Matrix Spike Sample (PDS)	or MSD fail	Recovery ±20% of true value	Analyze serial dilution of sample
	Serial Dilution Test (L)		1:5 dilution of sample must agree within 10% with undiluted result	If MS, MSD, PDS, and serial dilution fail, quantitate sample by method of standard additions

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#### TABLE 1

#### QC REQUIREMENTS (CONTINUED)

Parameter/ Method	QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action				
	Instrument Detection Limit (IDL) Study	Quarterly.	uarterly. IDL < PQL 1)Repeat IDL study. 2)Raise PQL.					
7471B	Method Detection Limit (MDL) Study		x-806, "Method Detection Limi cudies and Verifications", curre	,				
	Limit of Detection (LOD) determination	Quarterly.	LOD = 2-3X MDL	Repeat LOD Determination.				

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#### TABLE 2

#### DoD QSM QC REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Demonstrate acceptable analytical capability	Prior to using any test method and at any time there is a significant change in instrument type, personnel, test method, or sample matrix.	QC acceptance criteria published by DoD, if available; otherwise, method-specified criteria.	Recalculate results; locate and fix problem, then rerun demonstration for those analytes that did not meet criteria.	NA.	This is a demonstration of analytical ability to generate acceptable precision and bias per the procedure in Appendix C. No analysis shall be allowed by analyst until successful demonstration of capability is complete.
LOD determination and verification	(Refer to current revision of SOP QA- 806)				
LOQ establishment and verification	(Refer to current revision of SOP QA- 806)				
Initial calibration (ICAL) for Mercury: minimum 5 standards and a calibration blank	Daily ICAL prior to sample analysis.	If more than one calibration standard is used, r ≥ 0.995.	Correct problem, then repeat ICAL.	Flagging criteria are not appropriate.	Problem must be corrected. No samples may be run until ICAL has passed.
Second source calibration verification (ICV)	Once after each ICAL, prior to beginning a sample run.	Value of second source for all analyte(s) within ± 10% of true value.	Correct problem and verify second source standard. Rerun ICV. If that fails, correct problem and repeat ICAL.	Flagging criteria are not appropriate.	Problem must be corrected. No samples may be run until calibration has been verified.
Continuing calibration verification (CCV)	After every 10 field samples and at the end of the analysis sequence.	CVAA: within ± 20% of true value.	Correct problem, rerun calibration verification. If that fails, then repeat ICAL. Reanalyze all samples since the last successful calibration verification.	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply Q-flag to all results for the specific analyte(s) in all samples since the last acceptable calibration verification.	Problem must be corrected. Results may not be reported without a valid CCV. Flagging is only appropriate in cases where the samples cannot be reanalyzed.

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#### TABLE 2

#### DoD QSM QC REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Method blank	One per preparatory batch.	No analytes detected > ½ RL and > 1/10 the amount measured in any sample or 1/10 the regulatory limit (whichever is greater). Blank result must not otherwise affect sample results. Contact Client if samples cannot be reprepped within hold time.  For negative blanks, absolute value < LOD.	Correct the problem. Report sample results that are <lod or="">10x the blank concentration. Reprepare and reanalyze the method blank and all associated samples with results &gt; LOD and &lt; 10x the contaminated blank result. Contact Client if samples cannot be reprepped within hold time.</lod>	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply B-flag to all results for the specific analyte(s) in all samples in the associated preparatory batch.	Problem must be corrected. Results may not be reported without a valid method blank. Flagging is only appropriate in cases where the samples cannot be reanalyzed.
Calibration blank	Before beginning a sample run, after every 10 samples, and at end of the analysis sequence.	No analytes detected > LOD. For negative blanks, absolute value < LOD.	Correct problem. Reprep and reanalyze calibration blank. All samples following the last acceptable calibration blank must be reanalyzed.	Apply B-flag to all results for specific analyte(s) in all samples associated with the blank.	
LCS containing all analytes to be reported	One per preparatory batch.	Water: Recovery must be within ± 20% of the true value Soil: Recovery must be within vendor supplied limits (varies by lot).	Correct problem, then reprep and reanalyze the LCS and all samples in the associated preparatory batch for failed analytes, if sufficient sample material is available.	If reanalysis cannot be performed, data must be qualified and explained in the case narrative. Apply Q-flag to specific analyte(s) in all samples in the associated preparatory batch.	Problem must be corrected. Results may not be reported without a valid LCS. Flagging is only appropriate in cases where the samples cannot be reanalyzed.
Matrix spike (MS)	One per preparatory batch per matrix (see Box D-7).	Recovery must be within ± 20% of the true value	Examine the project- specific DQOs. If the matrix spike falls outside of DoD criteria, additional quality control tests are required to evaluate matrix effects.	For the specific analyte(s) in the parent sample, apply J-flag if acceptance criteria are not met.	For matrix evaluation only. If MS results are outside the LCS limits, the data shall be evaluated to determine the source of difference and to determine if there is a matrix effect or analytical error.
Matrix spike duplicate (MSD) or sample duplicate	One per preparatory batch per matrix	MSD: Recovery must be within ± 20% of the true value.  MSD or sample duplicate: RPD ≤ 20% (between MS and MSD or sample and sample duplicate).	Examine the project- specific DQOs. Contact the client as to additional measures to be taken.	For the specific analyte(s) in the parent sample, apply J-flag if acceptance criteria are not met.	The data shall be evaluated to determine the source of difference.

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#### TABLE 2

#### DoD QSM QC REQUIREMENTS

QC Check	Minimum Frequency	Acceptance Criteria	Corrective Action	Flagging Criteria	Comments
Method of standard additions (MSA)	When matrix interference is confirmed.	NA.	NA.	NA.	Document use of MSA in the case narrative.
Results reported between DL and LOQ	NA.	NA.	NA.	Apply J-flag to all results between DL and LOQ.	

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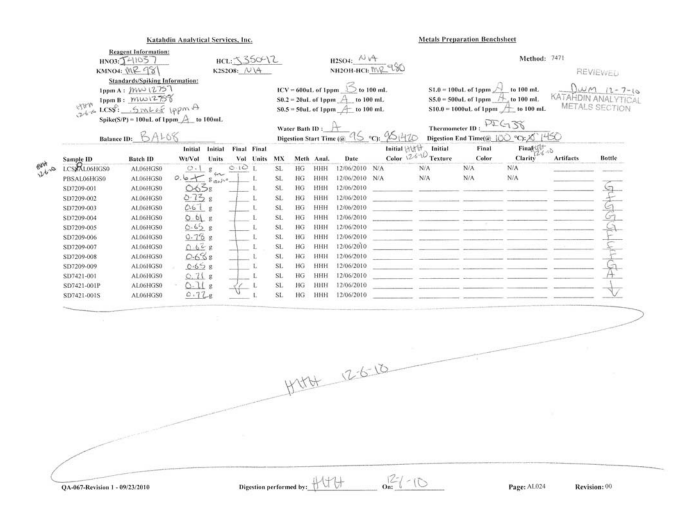
# TABLE 3 SUMMARY OF METHOD MODIFICATIONS

Topic	Katahdin SOP CA-611-09	USEPA Method 7471, current revision
Reagents	Stannous chloride dissolved in hydrochloric acid to prevent clogging of mercury analyzer, per instrument manufacturer's recommendation.	Stannous chloride dissolved/suspended in sulfuric acid.
Procedures	Sampling and gas stream switching performed automatically by mercury analyzer.	Sampling and gas stream switching performed manually by analyst.
QC – Calibration Verification	1)Known reference sample (ICV) analyzed daily. 2)Calibration verified after every 10 samples with CCV.	Nnown reference sample analyzed quarterly.     Calibration verified after every 20 samples.
QC - Calibration Blanks and Method Blanks	Acceptance Criterion: < PQL	Acceptance criteria: Low enough not to interfere with data quality objectives, or <10% of PQL, or <10% of regulatory limit, or <10% of lowest associated sample

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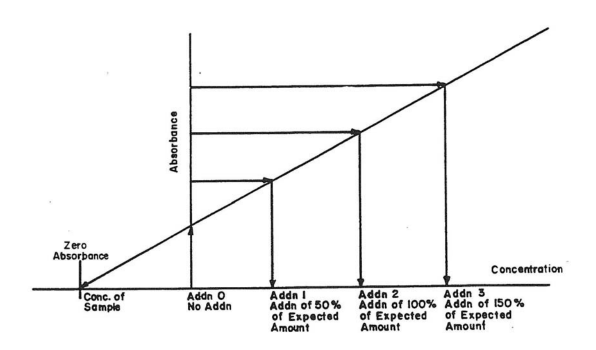
# FIGURE 1 EXAMPLE PAGE FROM MERCURY PREPARATION LOGBOOK



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FIGURE 2
STANDARD ADDITIONS PLOT



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#### FIGURE 3

#### EXAMPLE CERTIFICATE OF ANALYSIS FOR A SOLID REFERENCE MATERIAL



M51475

DataPacK<sup>TM</sup> Lot No. D051-540 **Trace Metals in Soil** 

Catalog No. 540

Certification

Method 3050 HNO3, H2O2, HCI	Total Concentration <sup>1</sup> (mg/Kg)	Certified Value <sup>2</sup> (mg/Kg)	Performance Acceptance Limits™ <sup>3</sup> (mg/Kg)
Parameter			1620 11100
aluminum	55600*	7870	4630 - 11100
antimony	160	70.5	D.L 149
arsenic	316	289	234 - 344
barium	869	211	174 - 247
beryllium	60.9	54.4	45.2 - 63.6
boron	129	91.3	58.8 - 124
cadmium	114	101	82.9 - 119
calcium	9750*	3680	2970 - 4390
chromium	249	224	180 - 268
cobalt	113	101	82.7 - 119
copper	94.9	88.0	73.3 - 103
iron	24400*	15700	6610 - 24900
lead	184	158	129 - 187
magnesium	3780*	2260	1760 - 2750
manganese	703	420	343 - 497
mercury	5.32	5.18	3.42 - 6.87
molybdenum	80.2	69.6	55.5 - 83.7
nickel	137	120	99.1 - 141
potassium	33000*	3000	2200 - 3800
selenium	146		101 - 159
silver	127	130	68.9 - 139
sodium	15600*	104	692 - 1470
strontium		1080	90.5 - 135
thallium	326	113	72.8 - 115
tin	106	94.0	104 - 194
titanium	175	149	116 - 453
vanadium	3100*	284	85.1 - 137
	151	111	215 - 329
zinc	311	272	213 - 329

	Total	Certified	Performance
Method 3050 HNO3, H2O2	Concentration 1	Value 2	Acceptance Limits <sup>™ 3</sup>
	mg/Kg	mg/Kg	mg/Kg
Parameter	5,5	mg/ reg	
aluminum	55600*	7380	4440 - 10300
antimony	160	75.2	D.L 198
arsenic	316	284	225 - 343
barium	869	217	177 - 257
beryllium	60.9	53.6	42.7 - 64.5
boron	129	89.5	58.9 - 120
cadmium	114	103	83.6 - 122
calcium	9750*	3540	2800 - 4270
chromium	249	224	172 - 275
cobalt	113	101	82.0 - 120
copper	94.9	85.5	70.4 - 100
iron	24400*	12500	5480 - 19500
lead	184	162	132 - 192
magnesium	3780*	2160	1650 - 2670
manganese	703	415	330 - 500
mercury	5.32	5.18	3.42 - 6.87
molybdenum	80.2	68.8	52.7 - 84.9
nickel	137	119	98.5 - 140
potassium	33000*	2840	2160 - 3520
selenium	146	135	104 - 166
silver	127	107	49.8 - 164
sodium	15600*	1010	709 - 1310
strontium	326	111	89.0 - 133
thallium	106	99.3	76.8 - 122
tin	175	148	70.6 - 225
titanium	3100*	283	104 - 463
vanadium	151	104	70.5 - 138
zinc	311	275	222 - 328
	011	2/3	

# ADDENDUM SOP NO CHANGE FORM

# KATAHDIN ANALYTICAL SERVICES, INC. SOP "REVIEW WITH NO CHANGES" FORM

Name of Person Reviewing SOP: Nicholas 11110	742011
Review Date: 1/28/13	
SOP Number: CA510-07	
SOP Title: TCLP for inorganic and non-vola	tile organic analytes
THE ABOVE REFERENCED SOP HAS BEEN REVIEWED ANALYST OR SUPERVISOR. NO CHANGES ARE REQU	BY A QUALIFIED AND TRAINED IRED TO THE SOP AT THIS TIME
Department Supervisor Signature:	Date:
Atherine	02/26/13
QAO Signature:	Date:
Leseis Dimond	030413



#### SCOPE OF ACCREDITATION TO ISO/IEC 17025:2005

#### TESTAMERICA LABORATORIES WEST SACRAMENTO

880 Riverside Parkway West Sacramento, CA 95605 Douglas Weir Phone: 916 3744 389

#### **ENVIRONMENTAL**

Valid To: January 31, 2014 Certificate Number: 2928.01

In recognition of the successful completion of the A2LA evaluation process, (including an assessment of the laboratory's compliance with ISO IEC 17025:2005, the 2003 NELAC Chapter 5 Standard, and the requirements of the DoD Environmental Laboratory Accreditation Program (DoD ELAP) as detailed in version 4.2 of the DoD Quality Systems Manual for Environmental Laboratories) accreditation is granted to this laboratory to perform recognized EPA methods using the following testing technologies and in the analyte categories identified below:

#### **Testing Technologies**

Inductively Coupled Plasma (ICP), ICP-Mass Spectroscopy, Atomic Absorption Spectroscopy (flame), Gas Chromatography(GC), GC- Mass Spectroscopy, High Resolution Gas Chromatography/High Resolution Mass Spectroscopy, Liquid Chromatography(LC), LC- Mass Spectroscopy, Ion Chromatography, Spectrophotometry, Misc.- Electronic Probes

Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
<u>Metals</u>				
Aluminum	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Antimony	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Arsenic	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Barium	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Beryllium	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Cadmium	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Calcium	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Chromium (Total)	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Chromium (Hexavalent)	EPA 7196A	EPA 7196A		EPA 3005A/3010A/3050A
Cobalt	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Copper	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Iron	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Lead	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Magnesium	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A

Peter Mbryer

Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	Associated Prep Methods
Manganese	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Mercury	EPA 7470A	EPA 7471A		EPA 3005A/3010A/3050A
Molybdenum	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Nickel	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Potassium	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Selenium	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Silver	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Sodium	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Thallium	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Vanadium	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Zinc	EPA 6010B/6020	EPA 6010B/6020		EPA 3005A/3010A/3050A
Nutrients				
Nitrate	EPA 353.2/9056A/300.0	EPA 353.2/9056A/300.0		
Nitrate-nitrite	EPA 353.2/	EPA 353.2		
	SM4500-NO3 F			
Nitrite	EPA 353.2/9056A/300.0	EPA 353.2/9056A/300.0		
Orthophosphate	EPA 9056A/300.0	EPA 9056A/300.0		
Wet Chemistry				
0.1 1.0	EDA 16644 (2072)	ED 4 1 ( ( 4 4 /0071		
Oil and Grease	EPA 1664A/9070	EPA 1664A/9071		
Nitrocellulose	WS-WC-0050	WS-WC-0050		
Perchlorate	EPA 6850	EPA 6850		
Chloride	EPA 9056A/300.0	EPA 9056A/300.0		
Fluoride	EPA 9056A/300.0	EPA 9056A/300.0		
Sulfate	EPA 9056A/300.0	EPA 9056A/300.0		
II 1 W				
<u>Hazardous Waste</u> Characteristics				
Characteristics				
TCLP Extractables		EPA 1311		
TCLP Inorganics		EPA 1311		
TCLI morganics		ELATIT		
Purgeable Organics				
(volatiles)				
(volatiles)				
Acetone	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Acrolein	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Acrylonitrile	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Allyl Chloride	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Benzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Bromochloromethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Bromodichloromethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Bromoform	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Bromomethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Carbon disulfide	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Carbon tetrachloride	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Chlorobenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
CHIOLOGUZCHE	LI A 0200D/0200C	LIA 6200D/6200C		LI 1 30301/3030D/3033/3033A

Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
Chloroethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
2-Chloroethyl vinyl ether	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Chloroform	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Chloromethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Chloroprene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Cyclohexane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1-Chlorocyclohexane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Dibromochloromethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,2-Dibromo-3-	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
chloropropane				
1,2-Dibromoethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Dibromomethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,2-Dichlorobenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,3-Dichlorobenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,4-Dichlorobenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
T-1,4-Dichloro-2-Butene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Dichlorodifluoromethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,1-Dichloroethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,2-Dichloroethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,1-Dichloroethene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
trans-1,2-Dichloroethene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
cis-1,2-Dichloroethene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,2-Dichloropropane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,3-Dichloropropane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
2,2-Dichloropropane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,1-Dichloropropene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
cis-1,3-Dichloropropene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
trans-1,3-Dichloropropene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,4-Dioxane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Ethylbenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Ethylmethacrylate	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Hexachlorobutadiene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Hexane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
2-Hexanone (MBK)	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Iodomethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Isobutanol	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Isopropyl Ether	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Methacrylonitrile	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Methyl tert-butyl ether	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
(MTBE)	LI A 6200D/6200C	El A 8200B/8200C		LI A 3030A/3030B/3033/3033A
Methylene chloride	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Methyl ethyl ketone	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Methyl Methacrylate	EPA 8260B/8260C	EPA 8260B/8260C		
4-Methyl-2-pentanone	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
(MIBK)	LI A 6200D/6200C	El A 8200B/8200C		LI A 3030A/3030B/3033/3033A
Naphthalene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Propionitrile	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,1,1,2-Tetrachloroethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,1,2,2-Tetrachloroethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Tetrachloroethene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A  EPA 5030A/5030B/5035/5035A
Tetrahydrofuran	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A  EPA 5030A/5030B/5035/5035A
Tenanyuroruran	E1 A 0200D/0200C	EI A 0200D/0200C		LI A JUJUA/JUJUD/JUJJ/JUJJA

Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
Toluene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,2,3-Trichlorobenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,2,4-Trichlorobenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,1,1-Trichloroethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,1,2-Trichloroethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,1-2-Trichloro-1,2-2-	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
trifluorethane				
Trichloroethene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Trichlorofluoromethane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,2,3-Trichloropropane	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Vinyl acetate	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Vinyl chloride	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
m & p xylene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
o-xylene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Xylenes, Total	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
tert-amyl methyl ether	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
(TAME)				
tert-butyl alcohol (TBA)	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Ethyl tert-butyl ether (ETBE)	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Bromobenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
n-Butylbenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
sec-Butylbenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
tert-Butylbenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
2-Chlorotoluene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
4-Chlorotoluene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Isopropylbenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
n-Propylbenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Styrene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,2,4-Trimethylbenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
1,3,5-Trimethylbenzene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Oxygenates	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Gasoline Range Organics (GRO)	EPA 8260B/AK101	EPA 8260B/AK101		EPA 5030A/5030B/5035/5035A
p-Isopropyltoluene	EPA 8260B/8260C	EPA 8260B/8260C		EPA 5030A/5030B/5035/5035A
Extractable Organics				
(semivolatiles)				
(Seriii voluciies)				
Acenaphthene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
rechapithene	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
	1415 000/ 445 1415 0000	Wis doo, We wis door	0006	Air: 3542/TO-13A
Acenaphthylene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
T S	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
Acetophenone	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
2-Acetylaminofluorene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
4-Aminobiphenyl	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Aniline	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A

Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
Anthracene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
7,12-	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
Dimethylbenz(a)anthracene				3550B/3550C/3580A
Aramite	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
		2111 02 / 0 0 / 02 / 02		3550B/3550C/3580A
Benzaldehyde	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
Benzurden y de	E111 027 0 C/ 027 0 B	2111 027 00/027 02		3550B/3550C/3580A
Dibenze(a,j)acridine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
Dioenze(a,j)acriume	E1 A 82/0C/82/0D	E1 A 82/0C/82/0D		3550B/3550C/3580A
Benzidine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
Benziume	EPA 82/0C/82/0D	EPA 82/0C/82/0D		
D () (1	ED A 0270 C (0270 D /W/G	ED 1 0270 C /0270 D /W/G	IIIG	3550B/3550C/3580A
Benzo(a)anthracene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
Benzo(b)fluoranthene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
Benzo(k)fluoranthene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
Benzo(j)fluoranthene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
Benzogjinuorummene	E111 0270 C7027 0B	E111 027 0C/027 0B		3550B/3550C/3580A
Benzo(g,h,i)perylene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
Denzo(g,n,1)peryrene	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
	1V15-000/ VV 5-1V15-0008	1V13-000/ W 3-1V13-0008	0006	
D ( . )	EDA 0270C/9270D/W/C	EDA 92700/9270D/N/G		Air: 3542/TO-13A
Benzo(a)pyrene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
Benzo(e)pyrene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
Benzoic acid	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
				Air: 3542/TO-13A
Benzyl alcohol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
J 3 2				3550B/3550C/3580A
Benzyl butyl phthalate	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
20112yi outyi piitiididte	2111 02/00/02/00	2171 02700/02700		3550B/3550C/3580A
2-sec-Butyl-4,6-	EPA 8270C/8270D	EPA 8270C/8270D	+	EPA 3500B/3500C/3510C/
	EFA 02/0C/02/0D	EFA 02/0C/02/0D		
dinitrophenol	EDA 9270G/9270D	EDA 9270C/9270D	1	3550B/3550C/3580A
Bis(2-chloroethoxy)	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
Methane	FD 1 0050 5/50===	TD 1 0050 G/0555	1	3550B/3550C/3580A
Bis(2-chloroethyl) ether	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Bis(2-chloroisopropyl) ether	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
			<u> </u>	3550B/3550C/3580A
Di(2-ethylhexyl) phthalate	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
, , , , , , , , , , , , , , , , , , ,				3550B/3550C/3580A
4-Bromophenyl phenyl ether	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
. Bromophenyi pilenyi etilei	211102700702700	2111 027 007 027 00		3550B/3550C/3580A
Carbazole	EPA 8270C/8270D	EPA 8270C/8270D	+	EPA 3500B/3500C/3510C/
Cardazore	ErA 62/0C/82/0D	EFA 02/UC/02/UD		
				3550B/3550C/3580A

Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
4-Chloroaniline	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
4-Chloro-3-methylphenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Hexacloropropene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
1 1				3550B/3550C/3580A
2-Chloronaphthalene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
2-Chlorophenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
2 cmerephener	2111 021 0 21 021 02	2111 02 / 0 0 / 02 / 02		3550B/3550C/3580A
4-Chlorophenyl phenyl ether	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
4-Chlorophenyi phenyi emer	E174 0270C/0270B	E171 0270C/0270B		3550B/3550C/3580A
Charrygono	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
Chrysene				
	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
D: 11 -	ED 1 0050 G/0050D	ED 1 0050 G/0050D	0006	Air: 3542/TO-13A
Diallate	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3510C/3550B/
				3580A
Dibenz(a,h)anthracene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
Dibenzofuran	EPA 8270C/8270D	EPA 8270C/8270D	WS-	EPA 3500B/3500C/3510C/
			MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
1,2-Dichlorobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
1,2 Biemoroochzene	E111 027 0 C/ 027 0 B	2111 027 067 027 08		3550B/3550C/3580A
1,3-Dichlorobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
1,3-Dichiolobelizelle	EFA 82/0C/82/0D	EFA 82/0C/82/0D		3550B/3550C/3580A
1 4 D' 111	EDA 9270C/9270D	EDA 9270C/9270D		
1,4-Dichlorobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
		77.1.04.70.7(04.70.7)		3550B/3550C/3580A
3,3'-Dichlorobenzidine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
2,4-Dichlorophenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
2,6-Dichlorophenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Diethyl Phthalate	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
•				3550B/3550C/3580A
2,4-Dimethylphenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
, J F				3550B/3550C/3580A
Dimethyl Phthalate	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
2 miletily i i milatave	211102700702702	2111 02 / 0 0 / 02 / 02		3550B/3550C/3580A
Di-n-butyl phthalate	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
Di-ii-outyi piitiiaiate	E1 A 82/0C/82/0D	E1 A 82/0C/82/0D		3550B/3550C/3580A
D'	EDA 9270C/9270D	EDA 9270C/9270D		
Di-n-octyl phthalate	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
465: 1	ED 1 0050 G/0050D	ED 1 0050 G/0050D		3550B/3550C/3580A
4,6-Dinitro-2-methylphenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
2,4-Dinitrophenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
			1	3550B/3550C/3580A
2,4-Dinitrotoluene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
2,6-Dinitrotoluene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
,				3550B/3550C/3580A
1,4-Dioxane	WS-MS-0011	WS-MS-0011		EPA 3500B/3500C/3510C/

Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	Associated Prep Methods
1,2-Diphenylhydrazine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
(as Azobenzene)				3550B/3550C/3580A
Famphur	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Fluoranthene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
Fluorene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
Hexachlorobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Hexachlorobutadiene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Hexachlorocyclopentadiene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Hexachloroethane	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Indeno(1,2,3-c,d)pyrene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
Isodrin	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Isophorone	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
•				3550B/3550C/3580A
Isosafrole	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Isosafrole #1	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Isosafrole #2	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Kepone	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
•				3550B/3550C/3580A
Dimethoate	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Methapyrilene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
13				3550B/3550C/3580A
Methyl methanesulfonate	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
Methyl Parathion	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
3				3550B/3550C/3580A
3-Methylcholanthrene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
<b>,</b>				3550B/3550C/3580A
2-Methylnaphthalene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
J	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
2-Methylphenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
J. F				3550B/3550C/3580A
3-Methylphenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
4-Methylphenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
tonij ipiionoi	211102100102100	2111 02 100 102 100		3550B/3550C/3580A
3&4-Methylphenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
Jack intermy iphonor	2111 02/00/02/00	L111 02 / 0C/02 / 0D		3550B/3550C/3580A

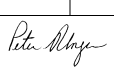


Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	Associated Prep Methods
Naphthalene	EPA 8270C/8270D/WS-	EPA 8270C/8270D/WS-	WS-	EPA 3500B/3500C/3510C/
•	MS-006/WS-MS-0008	MS-006/WS-MS-0008	MS-	3550B/3550C/3580A
			0006	Air: 3542/TO-13A
1,4-Naphthoquinone	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
1-Chloronaphthalene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
•				3550B/3550C/3580A
1-Methylnaphthalene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
3 1				3550B/3550C/3580A
1-Naphthylamine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
1 3				3550B/3550C/3580A
2-Naphthylamine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
2-Nitroaniline	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
	211102700702702	2111 027 0 07 027 02		3550B/3550C/3580A
3-Nitroaniline	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
3 Tittoumme	E111 0270C/0270B	E171 0270 C/0270D		3550B/3550C/3580A
4-Nitroaniline	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
4-1VIIIOammine	E174 0270C/0270B	E174 0270 C/0270D		3550B/3550C/3580A
4-Nitro-o-toluidine-1-oxide	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
4-1\lito-o-totuldilic-1-oxide	LI A 62/0C/62/0D	El A 62/0C/82/0D		3550B/3550C/3580A
5-Nitro-o-toluidine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
3-Milio-o-toluldine	EPA 82/0C/82/0D	EPA 82/0C/82/0D		
NI'a1	EDA 0270C/0270D	FDA 9270C/9270D		3550B/3550C/3580A
Nitrobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
100:4	ED 1 0270 C/0270 D	ED 4 0270 C/0270 D		3550B/3550C/3580A
1,2-Dinitrobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
100:1		FD 1 00 50 G/00 50 D		3550B/3550C/3580A
1,3-Dinitrobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
1,4-Dinitrobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
1,3,5-Trinitrobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
2-Nitrophenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
4-Nitrophenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
n-Nitrosodimethylamine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
n-Nitrosodi-n-propylamine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
n-Nitrosodiphenylamine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
n-Nitrosodiethylamine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
-				3550B/3550C/3580A
n-Nitroso-di-n-butylamine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
,				3550B/3550C/3580A
n-Nitrosomethylethylamine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
J J				3550B/3550C/3580A
n-Nitrosomorpholine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
1				3550B/3550C/3580A
n-Nitrosopyrrolidine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
222 F.J 2141112				3550B/3550C/3580A
	-		+	
n-Nitrosopiperidine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/

Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
Parathion	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
p-Chorobenzilate	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
p-Dimthylaminoazobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Pentachlorophenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Pentaclorobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Pentacloronitrobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Pentacloroethane	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Phenacetin	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Phenanthrene	EPA 8270C/8270D/WS-MS-006/WS-MS-0008	EPA 8270C/8270D/WS- MS-006/WS-MS-0008	WS- MS- 0006	EPA 3500B/3500C/3510C/ 3550B/3550C/3580A Air: 3542/TO-13A
a,a-Dimethylphenethlamine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Phenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Biphenyl	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Diphenylamine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
p-Phenylenediamine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
2-Picoline	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Phorate	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Promamide	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Pyrene	EPA 8270C/8270D/WS-MS-006/WS-MS-0008	EPA 8270C/8270D/WS- MS-006/WS-MS-0008	WS- MS- 0006	EPA 3500B/3500C/3510C/ 3550B/3550C/3580A Air: 3542/TO-13A
Pyridine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Safrole	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Sulfotepp	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Disulfotone	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Ethylmethanesulfonate	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
Thionazin	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
1,2,4,5-Tetrachlorobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A
1,2,4-Trichlorobenzene	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/ 3550B/3550C/3580A



Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
2,4,5-Trichlorophenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
_				3550B/3550C/3580A
2,3,4,6-Tetrachlorophenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
_				3550B/3550C/3580A
2,3,5,6-Tetrachlorophenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
•				3550B/3550C/3580A
2,4,6-Trichlorophenol	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
1				3550B/3550C/3580A
Diesel Range Organics	EPA	EPA		EPA 3500B/3500C/3510C/
(DRO)	8015B/8015C/AK102	8015B/8015C/AK102		3550B/3550C/3580A
Residual Range Organics	AK103	AK103		EPA 3500B/3500C/3510C/
				3550B/3550C/3580A
o,o,o-TEPT	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
2,2,2				3550B/3550C/3580A
o-Toluidine	EPA 8270C/8270D	EPA 8270C/8270D		EPA 3500B/3500C/3510C/
o Totalanie	2111 027 037 027 03	E111 027 007 027 0B		3550B/3550C/3580A
				3330B/3330C/3300/1
Dioxins				
DIOVIIIO	+			
2270 T CDD	ED 4 0200 4 /0200 D /0200	ED 1 0200 1 /0200		
2,3,7,8-TeCDD	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
1 2 2 7 0 P CDD	/8290A/1613B	/8290A/1613B		
1,2,3,7,8-PeCDD	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
1,2,3,4,7,8-HxCDD	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
1,2,3,6,7,8-HxCDD	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
1,2,3,7,8,9-HxCDD	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
1,2,3,4,6,7,8-HpCDD	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
OCDD	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
2,3,7,8-TeCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
1,2,3,7,8-PeCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
2,3,4,7,8-PeCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
1,2,3,4,7,8-HxCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
1,2,3,6,7,8-HxCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
1,2,3,7,8,9-HxCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
2,3,4,6,7,8-HxCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
1,2,3,4,6,7,8-HpCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
1,2,3,4,7,8,9-HpCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
, , , , , , , <b>r</b> -	/8290A/1613B	/8290A/1613B		
OCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
~ <del>~ ~ ~</del>	/8290A/1613B	/8290A/1613B		
Total TCDD	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
10(01100)				
	/8290A/1613B	/8290A/1613B		



Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	Associated Prep Methods
Total PeCDD	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
Total HxCDD	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
Total HeptaCDD	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
-	/8290A/1613B	/8290A/1613B		
Total TCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
Total PeCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
Total HxCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
Total HpCDF	EPA 8280A/8280B/8290	EPA 8280A/8280B/8290		
	/8290A/1613B	/8290A/1613B		
Chemical Warfare				
Degradates				
<u>Degrauates</u>			1	
1,4-Dithiane	WS-MS-0003	WS-MS-0003		
Benzothiazole	WS-MS-0003	WS-MS-0003		
p-Chlorophenyl	WS-MS-0003	WS-MS-0003		
methylsulfide	WB WB 0003	WB WB 0003		
p-Chlorophenyl	WS-MS-0003	WS-MS-0003		
methylsulfoxide	W 5 W 5 W 5 W 5 W 5 W 5 W 5 W 5 W 5 W 5	WE WE GOOD		
p-Chlorophenyl	WS-MS-0003	WS-MS-0003		
methylsulfone	, , , , , , , , , , , , , , , , , , ,	\\ \bar{\bar{\bar{\bar{\bar{\bar{\bar{		
Chloropicrin	WS-MS-0003	WS-MS-0003		
Acetophenone	WS-MS-0003	WS-MS-0003		
2-Chloroacetophenone	WS-MS-0003	WS-MS-0003	†	
1,4-Oxathiane	WS-MS-0003	WS-MS-0003		
Dimethyl Disulfide	WS-MS-0003	WS-MS-0003		
	WS-LC-0004	WS-IVIS-0003	+	
Diisopropylmethylphosphate (DIMP)	W S-LC-0004	WS-LC-0004		
Dimethylmethylphosphonate	WS-LC-0004	WS-LC-0004		
(DMMP)				
Ethyl methylphosphonic acid	WS-LC-0004	WS-LC-0004		
(EMPA)				
Isopropyl methylphosphonic acid (IMPA)	WS-LC-0004	WS-LC-0004		
Methylphosphonic acid	WS-LC-0004	WS-LC-0004		
(MPA)		,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,		
Thiodiglycol (2,2'-	WS-LC-0004	WS-LC-0004		
Thiodiethanol) (TDG)	We have the	112 20 000.		
, , ,				
Nitroaromatics				
2-Amino-4,6-dinitrotoluene	EPA 8330A/8330B	EPA 8330A/8330B		
4-Amino-2,6-dinitrotoluene	EPA 8330A/8330B	EPA 8330A/8330B		
3,5-Dinitroaniline	EPA 8330B	EPA 8330B		
1,3-Dinitrobenzene	EPA 8330A/8330B	EPA 8330A/8330B		
2,4-Dinitrotoluene	EPA 8330A/8330B	EPA 8330A/8330B	<b> </b>	
2,6-Dinitrotoluene	EPA 8330A/8330B	EPA 8330A/8330B		
Ethylene glycol dinitrate	EPA 8330B	EPA 8330B		
Emyrche grycor dimuate	LI A 0330D	L1 A 0330D		



Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
Glycerol trinitrate	EPA 8330B	EPA 8330B		
(Nitroglycerin)				
Hexahydro-1,3,5-trinitro-	EPA 8330A/8330B	EPA 8330A/8330B		
1,3,5-triazine (Hexogen)				
Methyl-2,4,6-	EPA 8330A/8330B	EPA 8330A/8330B		
trinitrophenylnitramine	FR + 0220 + /0220F	ED 1 0000 1 (0000 D		
Nitrobenzene	EPA 8330A/8330B	EPA 8330A/8330B		
2-Nitrotoluene (o-	EPA 8330A/8330B	EPA 8330A/8330B		
Nitrotoluene)	ED 1 0220 1 /0220 D	ED 1 0220 1 /0220 D		
3-Nitrotoluene (m-	EPA 8330A/8330B	EPA 8330A/8330B		
Nitrotoluene)	EBA 02204/0220B	ED 4 0220 4 /0220 D		
4-Nitrotoluene (p-	EPA 8330A/8330B	EPA 8330A/8330B		
Nitrotoluene)	ED 4 0220 4 /0220 D	ED 1 0220 1 /0220 D		
Octahydro-1,3,5,7-	EPA 8330A/8330B	EPA 8330A/8330B		
tetranitro1,3,5,7-tetracine				
(Octogen)	ED 4 0220D	ED 1 0220D		
Picric acid	EPA 8330B	EPA 8330B		
Pentaerythritol tetranitrate	EPA 8330B	EPA 8330B		
1,3,5-Trinitrobenzene	EPA 8330A/8330B	EPA 8330A/8330B		
2,4,6-Trinitrotoluene	EPA 8330A/8330B			
, ,		EPA 8330A/8330B		
Nitroguanidine	WS-LC-0010	WS-LC-0010		
>T'.				
<u>Nitrosamines</u>				
3137', 1' d 1 '	WG 14G 0010	WG MG 0012		
N-Nitrosodimethylamine	WS-MS-0012	WS-MS-0012		
(NDMA)				
Perfluoro Compounds				
Permuoro Compounds				
Perfluorooctanoic acid	WS-LC-0020	WS-LC-0020		
Perfluorooctanoic acid  Perfluorooctane sulfonate				
Permuorooctane sunonate	WS-LC-0020	WS-LC-0020		
Pesticides/PCBs				
<u>Festicides/FCBs</u>				
Aldrin	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
Alum	1699	1699		3550B/3550C/3620B/3660A
a-BHC	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
а-ВПС	1699	1699		3550B/3550C/3620B/3660A
b-BHC	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
0-BHC	1699	1699		3550B/3550C/3620B/3660A
d-BHC	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
d-BHC	1699	1699		3550B/3550C/3620B/3660A
g-BHC (Lindane)	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
g-BHC (Ellidane)	1699	1699		3550B/3550C/3620B/3660A
a-Chlordane	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
a-Chiordalic	1699	1699		3550B/3550C/3620B/3660A
g-Chlordane	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
5 Cinordane	1699	1699		3550B/3550C/3620B/3660A
Oxy-Chlordane	EPA 1699	EPA 1699		EPA 3500B/3500C/3510C/
Ony Chicianic	L111 10//			3550B/3550C/3620B/3660A
				1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
4,4'-DDD	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/



Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	Associated Prep Methods
2,4'-DDD	EPA 1699	EPA 1699		EPA 3500B/3500C/3510C/
				3550B/3550C/3620B/3660A
4,4'-DDE	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
	1699	1699		3550B/3550C/3620B/3660A
2,4'-DDE	EPA 1699	EPA 1699		EPA 3500B/3500C/3510C/
				3550B/3550C/3620B/3660A
4,4'-DDT	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
	1699	1699		3550B/3550C/3620B/3660A
2,4'-DDT	EPA 1699	EPA 1699		EPA 3500B/3500C/3510C/
				3550B/3550C/3620B/3660A
Dieldrin	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
	1699	1699		3550B/3550C/3620B/3660A
Endosulfan I	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
	1699	1699		3550B/3550C/3620B/3660A
Endosulfan II	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
	1699	1699		3550B/3550C/3620B/3660A
Endosulfan sulfate	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
	1699	1699		3550B/3550C/3620B/3660A
Endrin	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
	1699	1699		3550B/3550C/3620B/3660A
Endrin aldehyde	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
	1699	1699		3550B/3550C/3620B/3660A
Endrin ketone	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
	1699	1699		3550B/3550C/3620B/3660A
Heptachlor	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
	1699	1699	1	3550B/3550C/3620B/3660A
Heptachlor epoxide	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
	1699	1699		3550B/3550C/3620B/3660A
Hexachlorobenzene	EPA 1699	EPA 1699		EPA 3500B/3500C/3510C/
36.1	ED 1 0001 1 0001 D /ED 1	ED 1 0001 1 10001 D IED 1		3550B/3550C/3620B/3660A
Methoxychlor	EPA 8081A/8081B/EPA	EPA 8081A/8081B/EPA		EPA 3500B/3500C/3510C/
G' M 11	1699	1699		3550B/3550C/3620B/3660A
Cis-Nonachlor	EPA 1699	EPA 1699		EPA 3500B/3500C/3510C/
T N 11	EDA 1600	EDA 1600	1	3550B/3550C/3620B/3660A
Trans-Nonachlor	EPA 1699	EPA 1699		EPA 3500B/3500C/3510C/
T 1	EDA 0001 A /0001 D /EDA	FDA 0001 A /0001 D /FDA		3550B/3550C/3620B/3660A
Toxaphene	EPA 8081A/8081B/EPA 1699	EPA 8081A/8081B/EPA 1699		EPA 3500B/3500C/3510C/
Mirex	EPA 1699	EPA 1699		3550B/3550C/3620B/3660A EPA 3500B/3500C/3510C/
Milex	EPA 1099	EPA 1099		
Chlordane (technical)	EPA 8081A/8081B	EDA 0001 A /0001D		3550B/3550C/3620B/3660A EPA 3500B/3500C/3510C/
Chiordane (technical)	EPA 8081A/8081B	EPA 8081A/8081B		3550B/3550C/3620B/3660A
			+	3330B/3330C/3020B/3000A
DCD (Arcalara)				
PCB (Aroclors)				
DCD 1016	ED 4 0002/0002 4	EDA 0002/0002A	1	EDA 2500D/2500G/2510G/
PCB-1016	EPA 8082/8082A	EPA 8082/8082A		EPA 3500B/3500C/3510C/
				3550B/3550C/3620B/3660A/
DCD 1221	ED A 9092/9092 A	EDA 0002/0002A	+	3620B/3665A
PCB-1221	EPA 8082/8082A	EPA 8082/8082A		EPA 3500B/3500C/3510C/
PCB-1232	EPA 8082/8082A	EPA 8082/8082A		3550B/3550C/3620B/3660A/
PCB-1242	EPA 8082/8082A	EPA 8082/8082A		3620B/3665A
PCB-1248	EPA 8082/8082A	EPA 8082/8082A		EPA 3500B/3500C/3510C/
PCB-1254	EPA 8082/8082A	EPA 8082/8082A		3550B/3550C/3620B/3660A/
PCB-1260	EPA 8082/8082A	EPA 8082/8082A		3620B/3665A

Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	Associated Prep Methods
PCB-1262	EPA 8082/8082A	EPA 8082/8082A		EPA 3500B/3500C/3510C/
PCB-1268	EPA 8082/8082A	EPA 8082/8082A		3550B/3550C/3620B/3660A/
PCB (congeners)				
PCB 1 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
TCD T (DZ)	mod	mod		
PCB 2 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
,	mod	mod		
PCB 3 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 4 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 5 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD ( (DZ)	mod	mod		
PCB 6 (BZ)	EPA 1668A mod/1668C mod	EPA 1668A mod/ 1668C mod		
PCB 7 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
TCD / (DZ)	mod	mod		
PCB 8 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
()	mod	mod		
PCB 9 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
,	mod	mod		
PCB 10 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 11 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 12 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 12 (D7)	mod = 1/1/600C	mod		
PCB 13 (BZ)	EPA 1668A mod/1668C mod	EPA 1668A mod/ 1668C mod		
PCB 14 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 CD 14 (DZ)	mod	mod		
PCB 15 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 CD 13 (BL)	mod	mod		
PCB 16 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
,	mod	mod		
PCB 17 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 18 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 19 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 20 (D7)	mod	mod		
PCB 20 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
PCB 21 (BZ)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
I CD 21 (DZ)	mod	mod		
PCB 22 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
- 32 <b></b> (BL)	mod	mod		
PCB 23 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
` '	mod	mod		
PCB 24 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 25 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		



Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
PCB 26 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 27 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 28 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 29 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 30 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
PCB 32 (BZ)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
	mod	mod		
PCB 31 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
I CD 31 (DZ)	mod	mod		
PCB 33 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 CD 33 (BL)	mod	mod		
PCB 34 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
10001(02)	mod	mod		
PCB 35 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 00 00 (00)	mod	mod		
PCB 36 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 05 00 (BL)	mod	mod		
PCB 37 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
()	mod	mod		
PCB 38 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 39 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
<b>\</b> /	mod	mod		
PCB 40 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 41 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 42 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 43 (BZ)	EPA 1668A mod/1668C			
DCD 44 (DZ)	mod	mod 1/1660G		
PCB 44 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 45 (DG)	mod	mod EPA 1668A mod/ 1668C		
PCB 45 (BZ)	EPA 1668A mod/1668C			
PCB 46 (BZ)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
rCD 40 (DZ)	mod mod/1008C	mod		
PCB 47 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C	<b>+</b>	
TCD 47 (DZ)	mod	mod		
PCB 48 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 49 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 50 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 51 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
<i>3 (22)</i>	mod	mod		
PCB 52 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		



Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
PCB 53 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 54 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 55 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 56 (BZ) PCB 57 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
PCB 58 (BZ)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
	mod	mod		
PCB 59 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 CD 39 (DZ)	mod	mod		
PCB 60 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
TCD 00 (BE)	mod	mod		
PCB 61 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
100 01 (02)	mod	mod		
PCB 62 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
102 02 (22)	mod	mod		
PCB 63 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 64 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
- 52 ( . ( <i>DL</i> )	mod	mod		
PCB 65 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 66 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 67 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 68 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 69 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 70 (BZ)	EPA 1668A mod/1668C			
DCD 71 (D7)	mod	mod 1/1660G		
PCB 71 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 70 (D7)	mod	mod		
PCB 72 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
PCB 73 (BZ)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
rcb /3 (bz)	mod mod/1008C	mod		
PCB 74 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C	<u> </u>	
PCB /4 (BZ)	mod	mod		
PCB 75 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 76 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 77 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 78 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
· · ()	mod	mod		
PCB 79 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		



Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
PCB 80 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 81 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 82 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 83 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 84 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
PCB 85 (BZ)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
	mod	mod		
PCR 86 (R7)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
PCB 86 (BZ)	mod	mod		
PCB 87 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
TCD 07 (BE)	mod	mod		
PCB 88 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 CB 00 (BZ)	mod	mod		
PCB 89 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
102 07 (32)	mod	mod		
PCB 90 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 91 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
- 32 /1 (32)	mod	mod		
PCB 92 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
/ - ()	mod	mod		
PCB 93 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
` '	mod	mod		
PCB 94 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 95 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 96 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 97 (BZ)	EPA 1668A mod/1668C			
7.C7 00 (7.5)	mod	mod		
PCB 98 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 00 (D7)	mod	mod		
PCB 99 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 100 (DZ)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
PCB 100 (BZ)				
PCB 101 (BZ)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
	mod	mod		
PCB 102 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C	<b>+</b>	
	mod	mod		
PCB 103 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 104 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 105 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 106 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		



Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
PCB 108 (BZ)/107 (IUPAC)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 109 (BZ)/108 (IUPAC)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 107 (BZ)/109 (IUPAC)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 110 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
7.57 111 (7.7)	mod	mod		
PCB 111 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
PCD 114 (DZ)	mod 1/1660G	mod 1/1660G		
PCB 112 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
PCD 112 (DZ)	mod	mod 1/1//00C		
PCB 113 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 114 (DZ)	mod	mod		
PCB 114 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 115 (D7)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
PCB 115 (BZ)				
DCD 116 (D7)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
PCB 116 (BZ)				
DCD 117 (D7)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
PCB 117 (BZ)				
PCB 118 (BZ)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
PCB 118 (BZ)	mod	mod mod 1008A mod/ 1008C		
PCB 119 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 CB 119 (BZ)	mod	mod		
PCB 120 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C	<u> </u>	
1 CB 120 (BZ)	mod	mod		
PCB 121 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 CD 121 (BZ)	mod	mod		
PCB 122 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
108 122 (82)	mod	mod		
PCB 123 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 124 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 125 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 126 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
, ,	mod	mod		
PCB 127 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 128 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 129 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 130 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 131 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 132 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 133 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod	<u> </u>	



Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	<b>Associated Prep Methods</b>
PCB 134 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 135 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 136 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 137 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 138 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DGD 440 (DG)	mod	mod		
PCB 139 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
P.CD 140 (P.Z)	mod	mod		
PCB 140 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 141 (DZ)	mod	mod		
PCB 141 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 142 (D7)	mod	mod		
PCB 142 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 142 (D7)	mod = 1/1/600	mod		
PCB 143 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 144 (D7)	mod = 1/1/69C	mod		
PCB 144 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 145 (D7)	mod = 1/1/09C	mod = 1/1/2000		
PCB 145 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 146 (D7)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
PCB 146 (BZ)				
PCB 147 (BZ)	mod EPA 1668A mod/1668C	mod EPA 1668A mod/ 1668C		
rCD 14/ (DZ)	mod mod/1008C	mod mod 1008A mod/ 1008C		
PCB 148 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
rCD 146 (DZ)	mod	mod		
PCB 149 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C	<b>+</b>	
1 CD 147 (DL)	mod	mod		
PCB 150 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		 
1 CD 130 (DZ)	mod	mod		
PCB 151 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 CD 131 (BZ)	mod	mod		
PCB 152 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 CD 132 (BZ)	mod	mod		
PCB 153 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
()	mod	mod		
PCB 154 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
( )	mod	mod		
PCB 155 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
( )	mod	mod		
PCB 156 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
,	mod	mod		
PCB 157 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
, ,	mod	mod		
PCB 158 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
,	mod	mod		
PCB 159 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
` /	mod	mod		
PCB 160 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
· /	mod	mod		



Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	Associated Prep Methods
PCB 161 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 162 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 163 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 164 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 165 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 166 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 167 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 168 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 169 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 170 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 171 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 172 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 173 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 174 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 175 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 176 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
D.CD 4.55 (D.C)	mod	mod		
PCB 177 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
D.CD 1=0 (D.D.)	mod	mod		
PCB 178 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
D. G. D. (D. C.)	mod	mod		
PCB 179 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
PGD 100 (DZ)	mod	mod 1/1660G		
PCB 180 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
PCD 101 (DZ)	mod	mod 1/1660G		
PCB 181 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 102 (DZ)	mod	mod		
PCB 182 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 102 (D7)	mod	mod		
PCB 183 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 194 (D7)	mod	mod		
PCB 184 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 195 (D7)	mod   EDA 1669 A mod/1669 C	mod EDA 1669A mod/1669C	+	
PCB 185 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 106 (D7)	mod   EDA 1669A mod/1669C	mod	1	
PCB 186 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 107 (DZ)	mod	mod	+	
PCB 187 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		



Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	Associated Prep Methods
PCB 188 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 189 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 190 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 191 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 192 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 193 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 194 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 195 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 196 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 197 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
P.G. 100 (P.G.)	mod	mod		
PCB 198 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 201 (BZ)/199 (IUPAC)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 100 (DZ) (200 (H ID A C)	mod	mod		
PCB 199 (BZ)/200 (IUPAC)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 200 (DZ)/201 (H IDAC)	mod	mod	1	
PCB 200 (BZ)/201 (IUPAC)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
DCD 202 (DZ)	mod	mod EPA 1668A mod/ 1668C		
PCB 202 (BZ)	EPA 1668A mod/1668C	mod 1008A mod/ 1008C		
PCB 203 (BZ)	mod EPA 1668A mod/1668C	EPA 1668A mod/ 1668C	l	
FCB 203 (BZ)	mod	mod		
PCB 204 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C	<b>+</b>	
1 CB 204 (BZ)	mod	mod		
PCB 205 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 CD 203 (BZ)	mod	mod mod		
PCB 206 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
1 CB 200 (BZ)	mod	mod		
PCB 207 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
102 207 (22)	mod	mod		
PCB 208 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
PCB 209 (BZ)	EPA 1668A mod/1668C	EPA 1668A mod/ 1668C		
	mod	mod		
Hormones, Steroids,				
Pharmaceuticals and				
Disinfection Byproducts			<u> </u>	
Acetominophen	WS-LC-0024			
Atenolol	WS-LC-0024			
Azithromycin	WS-LC-0024			
Carbadox	WS-LC-0024			
Carbamazepine	WS-LC-0024			



Parameter/Analyte	Non potable Water	Solid Hazardous Waste	Air	Associated Prep Methods
Clarithromycin	WS-LC-0024			
Diazepam	WS-LC-0024			
Digoxigenin	WS-LC-0024			
Digoxin	WS-LC-0024			
Diphenylhydramine	WS-LC-0024			
Fluoxetine	WS-LC-0024			
Flumequine	WS-LC-0024			
Gemfibrozil	WS-LC-0024			
Ibuprofen	WS-LC-0024			
Naproxen	WS-LC-0024			
Ormetoprim	WS-LC-0024			
Penicillin G	WS-LC-0024			
Sulfachloropyridazine	WS-LC-0024			
Sulfadiazine	WS-LC-0024			
Sulfamethizole	WS-LC-0024			
Sulfamethoxazole	WS-LC-0024			
Sulfathiazole	WS-LC-0024			
Thiabendazole	WS-LC-0024			
Trimethoprim	WS-LC-0024			
Tris (2-chloro-2-	WS-LC-0024			
propyl)phosphate (TCPP)				
Warfarin	WS-LC-0024			

Peter Mbryer



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In recognition of the successful completion of the A2LA evaluation process that includes an assessment of the laboratory's compliance with ISO/IEC 17025:2005, the 2003 NELAC Chapter 5 Standard, and the requirements of the Department of Defense Environmental Laboratory Accreditation Program (DoD ELAP) as detailed in version 4.2 of the DoD Quality System Manual for Environmental Laboratories (QSM); accreditation is granted to this laboratory to perform recognized EPA methods as defined on the associated A2LA Environmental Scope of Accreditation. This accreditation demonstrates technical competence for this defined scope and the operation of a laboratory quality management system (refer to joint ISO-ILAC-IAF Communiqué dated 8 January 2009).



Presented this 27<sup>th</sup> day of March 2012.

President & CEO

For the Accreditation Council Certificate Number 2928.01

Valid to January 31, 2014

Revised June 1, 2012

For the tests or types of tests to which this accreditation applies, please refer to the laboratory's Environmental Scope of Accreditation.



SOP No. WS-ID-0005, Rev. 7.5 Effective Date: 04/19/2013

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Title: Analysis of Samples for Polychlorinated Dioxins and Furans by HRGC/HRMS

[Methods 8290, 8290A & TO-9A]

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#### 1. SCOPE AND APPLICATION

- 1.1.1. This method provides procedures for the detection and quantitative measurement of 2,3,7,8-tetrachlorodibenzo-p-dioxin (2,3,7,8-TCDD), polychlorinated dibenzo-p-dioxins (tetra- through octachlorinated homologs; PCDDs), and polychlorinated dibenzofurans (tetra- through octachlorinated homologs; PCDFs) in a variety of environmental matrices at part-per-trillion (ppt) concentrations by SW 846 Method 8290 and 8290A. The analytical method calls for the use of high-resolution gas chromatography and high-resolution mass spectrometry (HRGC/HRMS) on purified sample extracts. An optional method for reporting the analytical results using a 2,3,7,8-TCDD toxicity equivalency factor (TEF) is also described. Table 1 lists the various sample types covered by this analytical protocol, the 2,3,7,8-TCDD-based method calibration limits and other pertinent information.
- 1.2. The sensitivity of this method is dependent upon the level of interferences within a given matrix.
- 1.3. This method is designed for use by analysts who are experienced with residue analysis and skilled in high-resolution gas chromatography/high resolution mass spectrometry (HRGC/HRMS).
- 1.4. Samples containing concentrations of specific congeners (PCDDs and PCDFs) that are greater than the calibration limit should be analyzed by a protocol designed for such concentrations, such as 8280A/B.
- 1.5. When undertaking projects for Department of Defense (DoD) the relevant criteria in QA Policy WS-PQA-021 "DoD QSM and AFCEE QAPP Implementation" must be checked and incorporated.

#### 2. SUMMARY OF METHOD

- 2.1. This procedure uses matrix-specific extraction, analyte-specific cleanup, and high-resolution capillary column gas chromatography/high resolution mass spectrometry (HRGC/HRMS) techniques. Sample preparation is addressed in WS-IDP-0005.
- 2.2. One to two  $\mu$ L of the concentrated extract are injected into an HRGC/HRMS system capable of performing selected ion monitoring at resolving powers of at least 10,000 (10 percent valley definition).
- 2.3. The identification of ten of the 2,3,7,8-substituted congeners (Table 3), for which a <sup>13</sup>C-labeled standard is included as a spiked compound, is based on their elution at their exact retention time (-1 to +3 seconds from the respective isotope dilution analyte or internal standard signal) and simultaneous detection of the two most abundant ions in

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the molecular ion region. All other identified PCDD/PCDF congeners are identified by their RRT's based on the daily CCV standard, and the simultaneous detection of the two most abundant ions in the molecular ion region. Confirmation is based on a comparison of the ratio of the integrated ion abundance of the molecular ion species to their theoretical abundance ratio.

2.4. Quantification of the individual congeners, total PCDDs and total PCDFs is achieved in conjunction with the establishment of a multipoint (five points) calibration curve for each homolog, during which each calibration solution is analyzed once.

#### 3. **DEFINITIONS**

- 3.1. Definitions of terms used in this SOP may be found in the glossary of the Quality Assurance Manual (QAM).
- 3.2. Data qualifiers are defined on each data report. Commonly used data qualifiers are defined in the QAM.
- 3.3. Polychlorinated dibenzo-p-dioxins (PCDDs) and polychlorinated dibenzo-furans (PCDFs): compounds (Figure 1) that contain from one to eight chlorine atoms. The seventeen 2,3,7,8-substituted PCDDs and PCDFs are shown in Table 3. The number of isomers at different chlorination levels is shown in Table 4
- 3.4. Homologous series: Defined as a group of chlorinated dibenzodioxins or dibenzofurans having a specific number of chlorine atoms.
- 3.5. Isomer: Chemical compounds that contain the same number of atoms of the same elements, but differ in structural arrangement and properties. For example, 1,2,3,4-TCDD and 2,3,7,8-TCDD are different structural isomers.
- 3.6. Congener: Any isomer of any homologous series.
- 3.7. Isotope Dilution Analyte: An isotope dilution analyte is a <sup>13</sup>C-labeled analog of a congener chosen from the compounds listed in Table 3. Isotope dilution analytes are added to all samples including method blanks and quality control samples before extraction, and they are used to quantitate the concentration of the analytes. Nine isotope dilution analytes are used in this method. There is one for each of the dioxin and furan homologs (except for OCDF) with the degree of chlorination ranging from four to eight. Additional isotope dilution analytes may be added to act as retention time references, but they are not used for quantitation.
- 3.8. Internal Standard: Two internal standards are used to determine the percent recoveries for the isotope dilution analytes. The <sup>13</sup>C-1,2,3,4-TCDD is used to measure the percent recoveries of the tetra- and pentachlorinated isotope dilution analytes while <sup>13</sup>C-1,2,3,7,8,9-HxCDD is used to determine the recovery of the hexa-, hepta- and

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octachlorinated isotope dilution analytes. <sup>13</sup>C-1,2,3,7,8,9-HxCDD also acts as a retention time reference for the unlabeled analog present in sample extracts. They are added to the final sample extract before HRGC/HRMS instrument analysis.

- 3.9. Estimated Detection Limit (EDL)/ Estimated Quantitation Limit (EQL): The sample specific estimated detection limit (EDL/EQL) is the concentration of a given analyte required to produce a signal with a peak height of at least 2.5 times the background noise level.
- 3.10. Estimated Maximum Possible Concentration (EMPC): The calculated concentration of a signal having the same retention time as a PCDD/PCDF congener, but which does not meet the other qualitative identification criteria defined in the method.

#### 4. INTERFERENCES

- 4.1. Solvents, reagents, glassware and other sample processing hardware may yield discrete artifacts or elevated baselines that may cause misinterpretation of the chromatographic data. All of these materials must be demonstrated to be free from interferents under the conditions of analysis by running laboratory method blanks. Analysts shall not use PVC gloves.
- 4.2. The use of high-purity reagents and solvents helps minimize interference problems. Purification of solvents by distillation in all-glass systems may be necessary.
- 4.3. Re-use of glassware is to be minimized to avoid the risk of contamination.
- 4.4. Interferents co-extracted from the sample will vary considerably from matrix to matrix. PCDDs and PCDFs are often associated with other interfering chlorinated substances such as polychlorinated biphenyls (PCBs), polychlorinated diphenyl ethers (PCDPEs), polychlorinated naphthalenes, and polychlorinated xanthenes that may be found at concentrations several orders of magnitude higher than the analytes of interest. Retention times of target analytes must be verified using reference standards. These values must correspond to the retention time windows established. While certain clean-up techniques are provided as part of this method, unique samples may require additional cleanup steps to achieve lower detection limits.
- 4.5. A high-resolution capillary column (60m DB-5) is used to resolve as many PCDD and PCDF isomers as possible. However, no single column is known to resolve all isomers. The DB-225 column is used for the quantitation of 2,3,7,8-TCDF when 2,3,7,8-TCDF on the DB-5 column is detected.

#### 5. SAFETY

Employees must abide by the policies and procedures in the Corporate Environmental Health and Safety Manual (CW-E-M-001), the Sacramento Addendum to the Corporate EH&S

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Manual (WS-PEHS-002) and this document. This procedure may involve hazardous material, operations and equipment. This SOP does not purport to address all of the safety problems associated with its use. It is the responsibility of the user of the method to follow appropriate safety, waste disposal and health practices under the assumption that all samples and reagents are potentially hazardous. Safety glasses, gloves, lab coats and closed-toed, nonabsorbent shoes are a minimum.

#### 5.1. Specific Safety Concerns or Requirements

- 5.1.1. The effluents of sample splitters for the gas chromatograph and roughing pumps on the HRGC/HRMS system should pass through either a column of activated charcoal or be bubbled through a trap containing oil or high-boiling alcohols.
- 5.1.2. Eye protection that satisfies ANSI Z87.1, laboratory coat, and chemically resistant gloves must be worn while samples, standards, solvents, and reagents are being handled. Latex and vinyl gloves provide no protection against most of the organic solvents used in this method. Nitrile or similar gloves must be used. Latex gloves may be used for methanol.
- 5.1.3. Exposure to chemicals must be maintained as low as reasonably achievable, therefore all samples must be opened, transferred and prepared in a fume hood. Solvent and waste containers will be kept closed unless transfers are being made.
- 5.1.4. Laboratory procedures such as repetitive use of pipets, repetitive transferring of extracts, and manipulation of filled separatory funnels and other glassware represent a significant potential for repetitive motion or other ergonomic injuries. Laboratory associates performing these procedures are in the best position to realize when they are at risk for these types of injuries. Whenever a situation is found in which an employee is performing the same repetitive motion, the employee shall immediately bring this to the attention of their supervisor, manager, or the EH&S staff. The task will be analyzed to determine a better means of accomplishing it.

#### 5.2. Primary Materials Used

The following is a list of the materials used in this method, which have a serious or significant hazard rating. **NOTE:** This list does not include all materials used in the method. The table contains a summary of the primary hazards listed in the MSDS for each of the materials listed in the table. A complete list of materials used in the method can be found in the reagents and materials section. Employees must review the information in the MSDS for each material before using it for the first time or when there are major changes to the MSDS.

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Material (1)	Hazards	Exposure Limit (2)	Signs and symptoms of exposure	
Acetone	Flammable	1000 ppm- TWA	Inhalation of vapors irritates the respiratory tract. May cause coughing, dizziness, dullness, and headache.	
Hexane	Flammable Irritant	500 ppm-TWA	Inhalation of vapors irritates the respiratory tract. Overexposure may cause lightheadedness, nausea, headache, and blurred vision. Vapors may cause irritation to the skin and eyes.	
Iso-octane	Flammable Irritant	None established	Inhalation of vapors may cause nausea, headache, dizziness, loss of consciousness, irritation to upper respiratory tract, pain in throat and nose, coughing, wheezing, shortness of breath.	
Methanol	Flammable Poison Irritant	200 ppm-TWA	A slight irritant to the mucous membranes. Toxic effects exerted upon nervous system, particularly the optic nerve. Symptoms of overexposure may include headache, drowsiness and dizziness. Methyl alcohol is a defatting agent and may cause skin to become dry and cracked. Skin absorption can occur; symptoms may parallel inhalation exposure. Irritant to the eyes.	
Methylene Chloride	Carcinogen Irritant	25 ppm-TWA 125 ppm-STEL	Causes irritation to respiratory tract. Has a strong narcotic effect with symptoms of mental confusion, light-headedness, fatigue, nausea, vomiting and headache. Causes irritation, redness and pain to the skin and eyes. Prolonged contact can cause burns. Liquid degreases the skin. May be absorbed through skin.	
Tetradecane	Irritant	None established	Inhalation of vapors may cause difficulty breathing, headache, intoxication and central nervous system damage.	
Toluene	Flammable Poison Irritant	200 ppm-TWA 300 ppm- Ceiling	Inhalation may cause irritation of the upper respiratory tract. Symptoms of overexposure may include fatigue, confusion, headache, dizziness and drowsiness. Peculiar skin sensations (e. g. pins and needles) or numbness may be produced. Causes severe eye and skin irritation with redness and pain. May be absorbed through the skin.	
		r to prevent violer		
2 – Exposure limit refers to the OSHA regulatory exposure limit.				

#### 6. EQUIPMENT AND SUPPLIES

- 6.1. Preventive and routine maintenance is described in the 'Schedule of Routine Maintenance' in the QAM.
- 6.2. High-Resolution Gas Chromatograph/High-Resolution Mass Spectrometer/Data System (HRGC/HRMS/DS).
  - 6.2.1. Capable of collecting, recording and storing MS data. The VG70 and Autospec Ultima systems utilize Opus version 3.6 software and the Autospec Premiere system utilizes MassLynx version 4.1 software.
  - 6.2.2. The GC must be equipped for temperature programming. All required accessories must be available, such as syringes, gases, and capillary columns. The GC injection port must be designed for capillary columns. The use of splitless injection techniques is recommended. The use of a moving needle

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injection port is also acceptable. When using the method described in this protocol, a 2- $\mu$ L injection volume is used consistently (i.e., the injection volumes for all extracts, blanks, calibration solutions and the performance check samples are 2  $\mu$ L). 1  $\mu$ L injections are allowed; however, laboratories are encouraged to remain consistent throughout the analyses by using the same injection volume at all times on a given HRGC/HRMS/DS.

- 6.2.3. Gas Chromatograph/Mass Spectrometer (GC/MS) Interface The GC/MS interface components should withstand 350° C. The interface must be designed so that the separation of 2,3,7,8-TCDD from the other TCDD isomers achieved in the gas chromatographic column is not appreciably degraded. Cold spots or active surfaces (adsorption sites) in the GC/MS interface can cause peak tailing and peak broadening. It is recommended that the GC column be fitted directly into the mass spectrometer ion source without being exposed to the ionizing electron beam. Graphite ferrules should be avoided in the injection port because they may adsorb the PCDDs and PCDFs. Vespel® or equivalent ferrules are recommended.
- 6.2.4. Mass Spectrometer The static resolving power of the instrument must be maintained at a minimum of 10,000 (10 percent valley). The mass spectrometer must be operated in a selected ion monitoring (SIM) mode with a total cycle time (including the voltage reset time) of one second or less.
- 6.2.5. Data System - A dedicated data system is employed to control the rapid multiple ion monitoring process and to acquire the data. Quantification data (peak areas or peak heights) and SIM traces (displays of intensities of each ion signal being monitored including the lock-mass ion as a function of time) must be acquired during the analyses and stored. Quantifications may be reported based upon computer-generated peak areas or upon measured peak heights (chart recording). The data system must be capable of acquiring data for a minimum of 10 ions in a single scan. It is also recommended to have a data system capable of switching to different sets of ions (descriptors) at specified times during an HRGC/HRMS acquisition. The data system should be able to provide hard copies of individual ion chromatograms for selected gas chromatographic time intervals. It should also be able to acquire massspectral peak profiles and provide hard copies of peak profiles to demonstrate the required resolving power. The data system should also permit the measurement of noise on the base line.

#### 6.3. GC Column

6.3.1. Due to poor separation of 2,3,7,8-TCDF from other TCDF isomers on the 60 m DB-5 column, a 30M DB-225 is used to quantitate 2,3,7,8-TCDF. This column is used when 2,3,7,8-TCDF is detected.

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6.3.2. In order to have an isomer-specific determination for 2,3,7,8-TCDD and to allow the detection of OCDD/OCDF within a reasonable time interval in one HRGC/HRMS analysis, the 60-m DB-5 fused-silica capillary column is recommended. At the beginning of each 12-hour period during which samples are analyzed and after tuning, acceptable compound separation on the GC column must be demonstrated through the analysis of a column performance check solution. Operating conditions known to produce acceptable results with the recommended column are shown in Table 7.

#### 7. REAGENTS AND STANDARDS

#### 7.1. Solvents

- 7.1.1. High-purity, distilled-in-glass or highest available purity: methylene chloride, hexane, methanol, tetradecane, isooctane, toluene, and acetone.
- 7.2. All calibration, daily isotope dilution analyte, daily clean up internal standards, and daily spiking solutions are stable for one year from preparation. After 1 year, solutions may be re-verified. The re-verified solution may be used for an additional year, or until there is evidence of compound degradation or concentration. The re-verification must be performed using an unexpired, not previously re-verified solution from a second lot or second vendor.
  - 7.2.1. Sealed ampules may be used until the manufacturer's expiration date is exceeded. If no expiration date is provided, then the expiration date will be 10 years from the date the ampule is opened. The solvent level should be monitored prior to each use to assure there has been no concentration of the standard over time.

#### 7.3. Calibration Solutions

- 7.3.1. High-Resolution Concentration Calibration Solutions (Table 5) Five tetradecane solutions containing unlabeled (totaling 17) and carbon-labeled (totaling 16) PCDDs and PCDFs at known concentrations are used to calibrate the instrument. The concentration ranges are homolog dependent, with the lowest values associated with the tetra chlorinated dioxins and furans (0.5 pg/μL) and the highest for the octachlorinated congeners (2000 pg/μL).
- 7.3.2. Individual isomers that make up the high-resolution concentration calibration solutions are obtained from commercial sources and prepared in the laboratory. These standards are traceable back to EPA-supplied standard solutions.
- 7.3.3. Store the calibration solutions in appropriate containers and at room temperature in the dark.

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7.3.4. Standards for method 8290A require storage at  $\leq 6^{\circ}$ C.

#### 7.4. GC Column Performance Check Solution

- 7.4.1. This solution contains the first and last eluting isomers for each homologous series from tetra- through hepta-chlorinated congeners. The solution also contains a series of other TCDD isomers for the purpose of documenting the chromatographic resolution. The <sup>13</sup>C-2,3,7,8-TCDD is also present. The laboratory is required to use tetradecane as the solvent and adjust the volume so that the final concentration does not exceed 100 pg/μL per congener. Table 8 summarizes the qualitative composition (minimum requirement) of this performance evaluation solution for the DB-5 column.
- 7.4.2. For the DB-225 column, the column performance check solution contains a series of TCDF isomers in addition to the 2,3,7,8-TCDF. The solution is injected and evaluated at the start of each analytical sequence on the DB-225 column to ensure that 2,3,7,8-TCDF is resolved from its closest eluting isomers with a baseline-to-valley ratio of  $\leq$  25%. Table 8 summarizes the qualitative composition (minimum requirement) of this performance evaluation solution on for the DB-225 column.
- 7.5. Field Surrogate Solution (air matrices)
  - 7.5.1. This solution contains one <sup>37</sup>Cl labeled analog (for Method TO-9/TO-9A) or one <sup>37</sup>Cl and four <sup>13</sup>C labeled analogs (for Methods 23 and/or 0023A) at the nominal concentration indicated in Table 2. It is used to assess sample collection and recovery procedures.
- 7.6. Sample Fortification Solution (Isotope dilution analyte)
  - 7.6.1. This isooctane (or toluene) solution contains the nine isotope dilution analytes at the nominal concentrations that are listed in Table 2. The solution contains at least one carbon-labeled standard for each homologous series, and it is used to measure the concentrations of the native substances. (Note that <sup>13</sup>C-OCDF is not present in the solution.)

#### 7.7. Internal Standard Solution

7.7.1. This tetradecane solution contains two internal standards (<sup>13</sup>C-1,2,3,4-TCDD and <sup>13</sup>C-1,2,3,7,8,9-HxCDD). An appropriate volume of this solution will be spiked into each sample extract before the final concentration step and HRGC/HRMS analysis.

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#### 8. SAMPLE COLLECTION, PRESERVATION AND STORAGE

- 8.1. The sample collection, shipping, handling, and chain-of-custody procedures are not described in this document. Sample collection personnel will, to the extent possible, homogenize samples in the field before filling the sample containers. This should minimize or eliminate the necessity for sample homogenization in the laboratory. The analyst should make a judgment, based on the appearance of the sample, regarding the necessity for additional mixing. If the sample is clearly non-homogeneous, the entire contents should be transferred to a glass or stainless steel pan for mixing with a stainless steel spoon or spatula before removal of a sample portion for analysis.
- 8.2. Grab and composite samples must be collected in glass containers.
- 8.3. Ambient air samples are collected on a Quartz Fiber Filter followed by a glass sleeve containing a polyurethane foam plug.
- 8.4. Samples from stationary sources are collected on glass or quartz fiber filters and XAD-2 Resin. (See WS-ID-0009 for sample preparation procedures).
- 8.5. Conventional sampling practices must be followed. Do not rinse the bottle with sample before collection. Sampling equipment must be free of potential sources of contamination.
- 8.6. With the exception of the fish tissues, which must be stored at  $20^{\circ}$ C, all samples should be stored at  $4^{\circ}$ C  $\pm$  2, extracted within 30 days and completely analyzed within 45 days of collection. The 30 day hold time is recommended. PCDDs and PCDFs have demonstrated stability for greater than one year.
- 8.7. All extracts must be stored capped, in the dark, at room temperature (approximately 21°C to 28°C). All extracts for method 8290A must be stored capped at  $\leq$  6°C.

#### 9. QUALITY CONTROL

9.1. One method blank (MB) must be extracted with every process batch of similar matrix, not to exceed twenty (20) samples. The method blank is an aliquot of laboratory matrix (reagent water, Ottawa sand, sodium sulfate, PUF, XAD, filter, etc.) processed in the same manner and at the same time as the associated samples. Corrective actions must be documented on a Non-Conformance memo, then implemented when target analytes are detected in the method blank above the reporting limit or when surrogate recoveries are outside control limits. Re-extraction of the blank, other batch QC, and the affected samples are required when the method blank is deemed unacceptable. The method blank contains a PUF plug, XAD, or filter prepared from the same batch as the field samples whenever possible for air samples.

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Certain programs, such as DOD, may require a more stringent evaluation of the method blank, for instance, that the blank not contain any analytes of interest at a concentration greater than ½ the lower calibration limit.

Note: Re-extraction of the blank, QC and affected samples for the air matrices (PUF, XAD, and filter) is not generally possible because the entire sample is consumed in the initial extraction.

- 9.1.1. The method blank must be spiked prior to extraction with the same amount of <sup>13</sup>C-labeled isotope dilution analytes as added to samples.
- 9.1.2. If method blank contamination is present, check solvents, reagents, fortification solutions, apparatus and glassware to locate and eliminate the source of contamination before any further samples are extracted and analyzed.
  - 9.1.2.1. OCDD is a ubiquitous laboratory contaminant. A method blank and the associated samples are deemed acceptable if the OCDD concentration is <5x the specified reporting limit. Flag data appropriately. The analyst is expected to investigate and eliminate potential sources of systematic contamination.
  - 9.1.2.2. If a target analyte is detected in the blank but the associated samples are ND (not detected), then the data may be reported, unless otherwise directed by the client. Note the action in the narrative.
  - 9.1.2.3. If a target analyte is detected in the blank, but the concentration of the contaminant in the samples >10x the blank concentration, then the data may be reported, unless otherwise directed by the client.

    Note the action in the narrative
- 9.1.3. If new batches of reagents or solvents contain interfering contaminants, purify or discard them.
- 9.2. A Laboratory Control Sample (LCS) must be extracted with every process batch of similar matrix, not to exceed twenty (20) samples. The LCS is an aliquot of laboratory matrix (e.g. water, Ottawa sand, sodium sulfate, PUF, XAD, etc.) spiked with analytes of known identity and concentration. The LCS must be processed in the same manner and at the same time as the associated samples. Corrective actions must be documented on a Non-Conformance memo, then implemented when recoveries of any spiked analyte is outside control limits provided on the LIMS or by the client. Reextraction of the blank, other batch QC and all associated samples are required if the LCS is deemed unacceptable. See policy WS-PQA-003 for specific acceptance criteria. When associated with PUF samples, the LCS should contain a PUF plug prepared from the same batch as the field samples whenever possible.

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Note: Re-extraction of the blank, QC and affected samples for the air matrices (PUF, XAD, and filter) is not generally possible because the entire sample is consumed in the initial extraction.

- 9.2.1. A LCS is deemed acceptable if control analytes are above upper control limits and the associated samples are ND, unless otherwise specified by the client. Note any actions in the narrative.
- 93 The assessment of matrix effects on method performance, as required by NELAP, is met in Method 8290 and 8290A, as in all isotope dilution techniques, with the use of isotopically labeled compounds. These isotopically labeled compounds are analogs of target analytes and are spiked into each sample. Therefore, matrix effects on method performance may be judged by the recovery of these analogs. Sample analysis acceptance is controlled by the performance of these analogs in each sample. A Matrix Spike/Matrix Spike Duplicate (MS/MSD or MS/SD) pair are extracted at the client's request only. Method 8290A does not address analysis of MS/MSD. An exception to this rule is a batch containing South Carolina samples for Method 8290. These batches must have an MS/MSD prepared. However, South Carolina requires Method 8290A after December 31, 2008. An MS/MSD pair are aliquots of a selected field sample spiked with analytes of known identity and concentration. When requested by the client, the MS/MSD pair shall be processed in the same manner and at the same time as the associated samples. Corrective actions must be documented on a Non-Conformance memo, then implemented when recoveries of any spike analyte is outside control limits provided on the LIMS or by the client. Re-extraction of the blank, the LCS, the selected field sample, and the MS/MSD may be required after evaluation and review. Matrix Spike/ Matrix Spike Duplicates are not generally applicable for air samples due to the difficulty in collecting identical or representative samples. An LCS/LCSD may be extracted to show precision of the extraction and analysis process.
  - 9.3.1. Matrix Spike (MS): A sample, which is spiked with a known amount of the matrix spike fortification solution prior to the extraction step. The recoveries of the matrix spike compounds are determined; they are used to estimate the effect of the sample matrix upon the analytical methodology.
  - 9.3.2. Matrix Spike Duplicate (MSD): A second portion of the same sample as used in the matrix spike analysis and which is treated like the matrix spike sample.
  - 9.3.3. Locate the sample for the MS and MSD analyses (the sample may be labeled "double volume").
  - 9.3.4. Add an appropriate volume of the matrix spike fortification solution, adjusting the fortification level as specified in Table 1, under IS Spiking Levels.
  - 9.3.5. Analyze the MS and MSD samples as described in Section 11.

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9.3.6. The results obtained from the MS and MSD samples (percent recovery and concentrations of 2,3,7,8-substituted PCDDs/PCDFs) should agree within 20 percent relative difference. Report all results and flag outliers.

9.3.7. Isotope dilution analyte recoveries are flagged if they are outside the recovery goals. Re-extraction of affected samples should be performed if signal-to-noise for any isotope dilution analyte is less than 10:1.

#### 9.4. Duplicates

- 9.4.1. Upon client request, duplicates may be processed. Locate the sample specified for duplicate analysis, and prepare and analyze a second 10-g soil or sediment sample portion or 1-L water sample, or an appropriate amount of the type of matrix under consideration. Duplicate samples are not generally applicable for air samples due to the difficulty in collecting identical or representative samples. A duplicate injection of a sample extract may be performed to display instrument precision.
  - 9.4.1.1. The results of the laboratory duplicates (percent recovery and concentrations of 2,3,7,8-substituted PCDD/PCDF compounds) should agree within 25 percent relative difference. Report all results and flag outliers.
- 9.4.2. Isotope dilution analyte recoveries are flagged if they are outside the recovery goals. Re-extraction of affected samples should be performed if signal-to-noise for any isotope dilution analyte is less than 10:1.

#### 9.5. Surrogate/Clean Up Internal Standard

A surrogate compound may be spiked into all air media samples prior to collection. For all other matrices, a clean up internal standard is spiked following extraction and just prior to cleanup, in order to monitor relative loss of isotope dilution analyte during both extraction and cleanup.

#### 9.6. Isotope Dilution Analytes

- 9.6.1. Isotope dilution analytes must be spiked into all samples, QC samples, and included in all calibrations.
- 9.6.2. For each sample and QC aliquot, calculate the percent recovery. The percent recovery should be between 40 percent and 135 percent for all nine isotope dilution analytes.
- 9.6.3. A low or high percent recovery for a blank does not require discarding the analytical data but it may indicate a potential problem with future analytical data. Isotope dilution analyte recoveries are flagged if they are outside the

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recovery goals. Re-extraction of affected samples should be performed if signal-to-noise for any isotope dilution analyte is less than 10:1.

- 9.7. Recommended Corrective Actions and Troubleshooting Steps
  - Verify satisfactory instrument performance.
  - If possible, verify that no error was made while weighing the sample portions.
  - Review the analytical procedures with the performing laboratory personnel.

#### 10. CALIBRATION

Calibration and Standardization requires a check of mass resolution (tuning), a check of chromatographic resolution, a verification of switching times (i.e. descriptors), and a calibration curve verification

- 10.1. For details of the calculations used to generate the regression equations, and how to use the factors generated by these equations, refer to SOP CA-Q-S-005 "Calibration Curves (General)".
- 10.2. Tuning (Mass Resolution Check)
  - 10.2.1. The mass spectrometer must be operated in the electron ionization mode. A static resolving power of at least 10,000 (10 percent valley definition) must be demonstrated at appropriate masses before any analysis is performed. Corrective actions must be implemented whenever the resolving power does not meet the requirement.
  - 10.2.2. Chromatography time for PCDDs and PCDFs exceeds the long-term mass stability of the mass spectrometer. Because the instrument is operated in the high-resolution mode, mass drifts of a few ppm (e.g., 5 ppm in mass) can have serious adverse effects on instrument performance. Therefore, a mass-drift correction is mandatory. To that effect, it is recommended to select a lockmass ion from the reference compound (PFK is recommended) used for tuning the mass spectrometer. The selection of the lock-mass ion is dependent on the masses of the ions monitored within each descriptor. Table 6 offers some suggestions for the lock-mass ions. However, an acceptable lock-mass ion at any mass between the lightest and heaviest ion in each descriptor can be used to monitor and correct mass drifts. The level of the reference compound (PFK) metered into the ion chamber during HRGC/HRMS analyses should be adjusted so that the amplitude of the most intense selected lock-mass ion signal (regardless of the descriptor number) does not exceed 10 percent of the full-scale deflection for a given set of detector parameters. Under those conditions, sensitivity changes that might occur during the analysis can be more effectively monitored.

*NOTE:* Excessive PFK (or any other reference substance) may cause noise problems and contamination of the ion source resulting in downtime for source cleaning.

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10.2.3. By using a PFK molecular leak, tune the instrument to meet minimum required resolving power of 10,000 (10 percent valley) at m/z 304.9824 (PFK) or any other reference signal close to m/z 303.9016 (from TCDF). Verify that the exact mass of m/z 380.9760 (PFK) is within 5 ppm of the required value. Note that the selection of the low- and high-mass ions must be such that they provide the largest voltage jump performed in any of the five mass descriptors (Table 6).

10.2.4. Documentation of the instrument resolving power must then be accomplished by recording the peak profile of the high-mass reference signal (m/z 380.9760). The minimum resolving power of 10,000 must be demonstrated on the high-mass ion while it is transmitted at a lower accelerating voltage than the low-mass reference ion, which is transmitted at full sensitivity. The format of the peak profile representation (Figure 3) must allow manual determination of the resolution, i.e., the horizontal axis must be a calibrated mass scale (amu or ppm per division). The result of the peak width measurement (performed at 5 percent of the maximum, which corresponds to the 10-percent valley definition) must appear on the hard copy and cannot exceed 100 ppm at m/z 380.9760 (or 0.038 amu at that particular mass).

#### 10.3. Performance Checks

- 10.3.1. At the beginning of each 12-hour period during which samples are to be analyzed, aliquots of the 1) GC column performance check solution and 2) high-resolution concentration calibration solution No. 4 (HRCC-4) shall be analyzed to demonstrate adequate GC resolution and sensitivity, response factor reproducibility, and mass range calibration, and to establish the PCDD/PCDF retention time windows. (Note: A HRCC-3 or HRCC-5 may be acquired to meet the requirement of #2 above. This is to provide documentation of consistency for varying concentration levels, and to meet NELAC requirements). A mass resolution check shall also be performed to demonstrate adequate mass resolution using an appropriate reference compound (PFK is recommended). If the required criteria are not met, remedial action must be taken before any samples are analyzed. The mass resolution check will be taken at the beginning and completion of an analytical sequence. An analytical sequence may contain one or more 12 hour periods.
  - 10.3.1.1 Method blanks or solvent blanks are used to demonstrate that the analytical system is free of contamination after the analysis of calibration standards or high level samples. The blank must demonstrate that the system has returned to appropriate background levels prior to continued analysis.
- 10.3.2. At a minimum, the ions listed in Table 6 for each of the five SIM descriptors

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must be monitored. Note that the PeCDF masses (M+2 & M+4) are also monitored in the first descriptor. This is because the first PeCDF isomer elutes closely to the final tetra isomer. The selection (Table 6) of the molecular ions M and M+2 for <sup>13</sup>C-HxCDF and <sup>13</sup>C-HpCDF rather than M+2 and M+4 (for consistency) is to eliminate, even under high-resolution mass spectrometric conditions, interferences occurring in these two ion channels for samples containing high levels of native HxCDDs and HpCDDs. It is important to maintain the same set of ions for both calibration and sample extract analyses. The recommended mass spectrometer tuning conditions are based on the groups of monitored ions shown in Table 6.

10.3.2.1. The GC column performance check mixture, high-resolution concentration calibration solutions, and the sample fortification solutions may be obtained from the EMSL-CIN. However, if not available from the EMSL-CIN, standards can be obtained from other sources, and solutions can be prepared in the laboratory. Concentrations of all solutions containing 2,3,7,8-substituted native PCDDs/PCDFs, must be verified by comparison with second-source standard solutions.

#### 10.4. Initial Calibration

Initial calibration is required before any samples are analyzed for PCDDs and PCDFs. Initial calibration is also required if any routine calibration (Section 10.5) does not meet the required criteria listed in Section 10.6.

- 10.4.1. Five high-resolution concentration calibration solutions, listed in Table 5, must be used for the initial calibration.
- 10.4.2. Tune the instrument with PFK.
- 10.4.3. Inject 1 or 2  $\mu$ L of the GC column performance check solution and acquire SIM mass spectral data as described earlier in Section 6.1.3. The total cycle time must be  $\leq$  1 second. This is analyzed prior to a calibration curve to set descriptor windows only and may not otherwise be documented. The laboratory must not analyze samples until it is demonstrated and documented that the criterion listed in Section 13.1 is met.
  - 10.4.3.1. Select the injection volume based upon the expected target analyte concentration, or expected matrix interferences.
  - 10.4.3.2. The same injection volume must be used for all samples, QC, and standards.
- 10.4.4. By using the same GC and mass spectrometer conditions that produced acceptable results with the column performance check solution, analyze a 1 or

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 $2-\mu L$  portion of each of the five concentration calibration solutions once with the following mass spectrometer operating parameter.

- 10.4.4.1. The total cycle time for data acquisition must be < 1 second. The total cycle time includes the sum of all dwell times and voltage reset times.
- 10.4.4.2. Acquire SIM data for all the ions listed in the five descriptors of Table 6.
- 10.4.4.3. The ratio of integrated ion current for the ions appearing in Table 9 (homologous series quantification ions) must be within the indicated control limits (set for each homologous series).
- 10.4.4.4. The ratio of integrated ion current for the ions belonging to the <sup>13</sup>C labeled isotope dilution analytes and internal standards must be within the control limits stipulated in Table 9.

NOTE: Section 10.4.3 requires that ion ratios be within the specified control limits simultaneously in one run. It is the laboratory's responsibility to take corrective action if the ion abundance ratios are outside the limits.

- 10.4.5. For each SICP and for each GC signal corresponding to the elution of a target analyte and of its labeled standards, the signal-to-noise ratio (S/N) must be better than or equal to 10. This measurement is suggested for any GC peak that has an apparent S/N of less than 5:1. The result of the calculation must appear on the SICP above the GC peak in question.
  - 10.4.5.1. Referring to Table 5, calculate the 17 relative response factors (RRF) for unlabeled target analytes [RRF(n); n=1 to 17] relative to their appropriate isotope dilution analytes (Table 5) and the nine RRFs for the labeled <sup>13</sup>C isotope dilution analytes [RRF(m); m=18 to 26] relative to the two internal standards according to the following formulae:

$$RRF(n) = \frac{A_x \times Q_{IDA}}{Q_x \times A_{IDA}}$$
  $RRF(m) = \frac{A_{IDA} \times Q_{IS}}{Q_{IDA} \times A_{IS}}$ 

Where

- A<sub>x</sub> = sum of the integrated ion abundances of the quantitation ions (Tables 6 and 5) for unlabeled PCDDs/PCDFs,
- $A_{IDA}$  = sum of the integrated ion abundances of the quantitation ions (Tables 6 and 5) for the labeled isotope dilution analytes,

 $A_{IS}$  = sum of the integrated ion abundances of the quantitation ions (Tables 6 and 10) for the labeled internal standards,

 $Q_{IDA}$  = quantity of the isotope dilution analyte injected (pg),

 $Q_{IS}$  = quantity of the internal standard injected (pg), and  $Q_x$  = quantity of the unlabeled PCDD/PCDF analyte injected (pg).

The RRF (n) and RRF (m) are dimensionless quantities; the units used to express  $Q_{IDA}$ ,  $Q_{IS}$ , and  $Q_X$  must be the same.

10.4.5.2. Calculate the RRF(n)s and their respective percent relative standard deviations (%RSD) for the five calibration solutions:

$$\overline{RRF}(n) = (\frac{1}{5}) \sum_{j=1}^{5} RRF_{j}(n)$$

Where n represents a particular PCDD/PCDF (2,3,7,8-substituted) congener (n = 1 to 17; Table 5), and j is the injection number (or calibration solution number; j = 1 to 5).

- 10.4.5.3. The relative response factors to be used for the determination of the concentration of total isomers in a homologous series are calculated as follows:
  - 10.4.5.3.1. For congeners that belong to a homologous series containing only one isomer (e.g., OCDD and OCDF) or only one 2,3,7,8-substituted isomer (Table 4; TCDD, PeCDD, HpCDD, and TCDF), the mean RRF used will be the same as the mean RRF determined in Section 10.3.5.2.

NOTE: The calibration solutions do not contain  $^{13}C$ -OCDF as an isotope dilution analyte. This is because a minimum resolving power of 12,000 is required to resolve the [M+6]+ ion of  $^{13}C$ -OCDF from the [M+2]+ ion of OCDD (and [M+4]+ from  $^{13}C$ -OCDF with [M]+ of OCDD). Therefore, the RRF for OCDF is calculated relative to  $^{13}C$ -OCDD.

10.4.5.3.2. For congeners that belong to a homologous series containing more than one 2,3,7,8-substituted isomer (Table 4), the mean RRF used for those homologous series will be the mean of the RRFs calculated for all individual 2,3,7,8-substituted congeners using the equation below:

$$\overline{RRF}(k) = (\frac{1}{t}) \sum_{n=1}^{t} RRF_n$$

Where:

t = total number of 2,3,7,8-substituted isomers present in the calibration solutions (Table 5) for each homologous series (e.g., two for PeCDF, four for HxCDF, three for HxCDD, two for HpCDF).

NOTE: Presumably, the HRGC/HRMS response factors of different isomers within a homologous series are different. However, this analytical protocol will make the assumption that the HRGC/HRMS responses of all isomers in a homologous series that do not have the 2,3,7,8-substitution patterns are the same as the responses of one or more of the 2,3,7,8-substituted isomer(s) in that homologous series.

10.4.5.4. Relative response factors [RRF(m)] to be used for the determination of the percent recoveries for the nine isotope dilution analytes are calculated as follows:

$$RRF(m) = \frac{A_{IDA}^{m} \times Q_{IS}}{Q_{IDA}^{m} \times A_{IS}}$$

$$\overline{RRF}(m) = (\frac{1}{5}) \sum_{j=1}^{5} RRF_{j}(m)$$

Where:

m = 18 to 26 (congener type) j = 1 to 5 (injection number),

 $A_{IDA}^{m}$  = sum of the integrated ion abundances of the

quantitation ions (Tables 6 and 10) for a given

isotope dilution analyte (m = 18 to 26),

 $A_{IDA}$  = sum of the integrated ion abundances of the

quantitation ions (Tables 6 and 10) for a given isotope dilution analyte (m = 18 to 26),

 $Q_{IDA} & Q_{IDA}^{m} =$  quantities of, respectively, the internal standard

(rs) and a particular isotope dilution analyte (m)

injected (pg),

RRF(m) = relative response factor of a particular isotope

dilution analyte (m) relative to an appropriate internal standard, as determined from one

injection, and

RRF(m) = calculated mean relative response factor of a particular isotope dilution analyte, as determined

from the five initial calibration injections (j).

10.5. Criteria for acceptable calibration

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The criteria listed below for acceptable calibration must be met before sample analysis is performed.

10.5.1. The percent relative standard deviations for the mean response factors [RRF(n) and RRF(m)] from the 17 unlabeled standards must be  $\leq$  20 percent, and those for the nine labeled reference compounds must be  $\leq$  30 percent.

Note: If Method 8290A criteria are required for the project then both the percent standard relative standard deviation for the mean response factors for the 17 unlabeled standards and the nine labeled reference compounds must be  $\leq 20$  percent.

- 10.5.2. The signal/noise ratio (S/N) for the GC signals present in every SICP (including the ones for the labeled standards) must be  $\geq 10$ .
- 10.5.3. The isotopic ratios (Table 9) must be within the specified control limits.

NOTE: If the criterion for acceptable calibration listed in Section 10.4.1 is met, the analyte-specific RRF can then be considered independent of the analyte quantity for the calibration concentration range. The mean RRFs will be used for all calculations until the routine calibration criteria (Section 10.6) are no longer met. At such time, new mean RRFs will be calculated from a new set of injections of the calibration solutions.

10.6. Routine Calibration (continuing calibration check)

Routine calibrations must be performed at the beginning of (following a successful tune and GC column performance check) and after a 12 hour period. The routine calibration initiates the 12 hour clock during which samples may be subsequently analyzed. The last sample in the sequence must be injected within 12 hours of the routine calibration, followed by the analysis of a closing calibration check. An acceptable closing calibration check standard may be used to initiate the next 12 hour analysis sequence when consecutive acquisition sequences occur. The ending mass resolution check shall be performed after the closing calibration check of an analysis acquisition sequence or after the final bracketing standard when consecutive 12 hour acquisition sequences are run.

- 10.6.1. Inject 1 or 2  $\mu$ L of the concentration calibration solution HRCC-4 containing 10 pg/ $\mu$ L of tetrachlorinated congeners, 50 pg/ $\mu$ L of penta-, hexa-, and heptachlorinated congeners, 100 pg/ $\mu$ L of octachlorinated congeners, and the respective isotope dilution analyte and internal standards (Table 5). By using the same HRGC/HRMS conditions as used in Sections 6.1.3 through 6.2, determine and document an acceptable calibration as provided in Section 10.6.
- 10.7. Criteria for Acceptable Routine Calibration

The following criteria must be met before further analysis is performed. If these criteria are not met, corrective action must be taken, including recalibration if needed.

- 10.7.1. The measured RRFs [RRF(n)] for the unlabeled standards obtained during the opening continuing calibration must be  $\pm$  20 percent of the mean values established during the initial calibration (Section 10.3.5.)
  - 10.7.1.1. The bracketing continuing calibration must be  $\pm$  20% of the average RRF calculated from the initial calibration.
    - 10.7.1.1.1 If the target compounds in the ending standard are less than or equal to  $\pm$  20% of the average RRF from the initial calibration, the RRFs of the initial calibration shall be used to quantitate the unlabeled isomers.
    - 10.7.1.1.2. If the target analytes are greater than  $\pm$  20% but less or equal to  $\pm$ 25% and the samples are non-detect, the data is acceptable and this anomaly is documented. If these isomers are greater than  $\pm$  20% but less or equal to  $\pm$ 25% and are positive, an average RRF of the initial and ending daily standard is calculated and used to quantitate the concentration of the affected congener, and the anomaly is documented.
    - 10.7.1.1.3. If the percent deviation of unlabeled compounds exceeds  $\pm$  25%, a new initial calibration is initiated within 2 hours following the analysis of the samples. Otherwise, reanalyze all sample extracts with positives for the failed target compounds.
- 10.7.2. The measured RRFs [RRF(m)] for the labeled standards obtained during the opening continuing calibration must be less than or equal to  $\pm$  30 percent of the mean values established during the initial calibration (Section 10.1.5).
  - 10.7.2.1. The bracketing continuing calibration must be  $\pm$  30% of the average RRF calculated from the initial calibration.
    - 10.7.2.1.1. If the labelled compounds in the ending standard are less than or equal to  $\pm 30\%$  of the average RRF from the initial calibration, the RRFs of the initial calibration shall be used to quantitate the labeled isomers.
    - 10.7.2.1.2. If the isotope dilution analyte analytes are greater than  $\pm$  30% but less or equal to  $\pm$ 35%, an average RRF of the initial and ending daily standards is calculated and used to quantitate the concentration of the affected congener.

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10.7.2.1.3. If the percent deviation of labeled compounds exceeds ± 35%, reanalyze samples if adversely impacted.

- 10.7.3. The ion-abundance ratios (Table 9) must be within the allowed control limits.
- 10.7.4. If either criteria in Sections 10.7.1 or 10.7.2 are not met, additional samples may not be analyzed. Sample data collected must be evaluated for usability. Narrate any reported data from the analytical sequence. If the ion-abundance ratio criterion is not satisfied, refer to the note in Section 10.4.3 for resolution.
- 10.7.5. If the above criteria (Section 10.7) cannot be satisfied, the entire initial calibration process (Section 10.4) must be repeated.

#### 11. PROCEDURE

#### 11.1. Procedural Variations

Procedural variations are allowed only if deemed necessary in the professional judgment of the supervisor to accommodate variation in sample matrix, radioactivity, chemistry, sample size, or other parameters. Any variation in procedure shall be completely documented using a Nonconformance memo and approved by a supervisor and QA/QC manager. If contractually required, the client will be notified. The Nonconformance memo will be filed in the project file.

Any deviations from this procedure identified after the work has been completed must be documented as a nonconformance, with a cause and corrective action described. A Nonconformance memo shall be used for this documentation.

#### 11.2. Sample Dilution Procedure – Simple Dilutions

Dilutions from 2X to 50X can be achieved without respiking the final extract. The calculation to determine the final extract concentration is as follows:

(Concentration of the original extract) x (amount of aliquot taken) x (volume of diluted extract) = final concentration of dilution.

Ex: 50X dilution of original 10 g/20  $\mu$ L sample (10 g/20  $\mu$ L) x (2  $\mu$ L aliquot + 98  $\mu$ L keeper) = 1 g/100  $\mu$ L FV

Record the final sample concentration on the extract label.

#### 11.3. Sample Dilution Procedure – Complex Dilutions

Complex dilution requiring respiking of IDA and IS: Dilutions greater than 50x must be done by diluting and respiking the extract with IDA's and IS. This procedure may require serial dilution to be performed. If this procedure is done, then the sample size must be adjusted to reflect the aliquot taken.

Ex. 100X dilution (original sample with 10 g/20 uL final volume)

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Take a 2  $\mu$ L aliquot (1/10 of original sample) and add 18  $\mu$ L of solvent keeper. Take a 2  $\mu$ L aliquot of the dilution (1/100 of the original sample), respike with 1 mL IDA and 20  $\mu$ L IS, reduced to 20  $\mu$ L FV.

Record the final sample concentration of the extract label.

#### 11.4. Analytical Procedures

- 11.4.1. Inject a 1 or 2  $\mu$ L aliquot of the extract into the GC, operated under the conditions previously used (Section 6.2) to produce acceptable results with the performance check solution.
- 11.4.2. Acquire SIM data according to Section 6.1.3. Use the same acquisition and mass spectrometer operating conditions previously used to determine the relative response factors (Section 10). Ions characteristic for polychlorinated diphenyl ethers are included in the descriptors listed in Table 6. Their presence is used to monitor their interference during the characterization of PCDFs.

#### 12. CALCULATIONS/DATA REDUCTION

#### 12.1. Identification Criteria

For a gas chromatographic peak to be identified as a PCDD or PCDF, it must meet all of the following criteria:

#### 12.1.1. Retention Times

- 12.1.1.1.For 2,3,7,8-substituted congeners, which have an isotopically labeled isotope dilution analyte or internal standard present in the sample extract, the retention time (at maximum peak height) of the sample components (i.e., the two ions used for quantitation purposes listed in Table 6) must be within -1 and +3 seconds of the retention time of the peak for the isotopically labeled isotope dilution analyte or internal standard at m/z corresponding to the first characteristic ion (of the set of two; Table 6) to obtain a positive identification of these nine 2,3,7,8-substituted PCDDs/PCDFs and OCDD.
- 12.1.1.2. For 2,3,7,8-substituted compounds that do not have an isotopically labeled isotope dilution analyte present in the sample extract, the relative retention time (relative to the appropriate isotope dilution analyte) must fall within 0.005 relative retention time units of the relative retention times measured in the daily routine calibration. Identification of OCDF is based on its retention time relative to <sup>13</sup>C-OCDD as determined from the daily routine calibration results.

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- 12.1.1.3. For non-2,3,7,8-substituted compounds (tetra through octa; totaling 119 congeners), the retention time must be within the corresponding homologous retention time windows established by analyzing the column performance check solution.
- 12.1.1.4. The ion current responses for both ions used for quantitative purposes (e.g., for TCDDs: m/z 319.8965 and 321.8936) must reach a maximum simultaneously (± 2 seconds).
- 12.1.1.5. The ion current responses for both ions used for the labeled standards (e.g., for <sup>13</sup>C-TCDD: m/z 331.9368 and m/z 333.9339) must reach a maximum simultaneously (± 2 seconds).
- 12.1.2. Ion Abundance Ratios

The integrated ion current for the two ions used for quantitation purposes must have a ratio between the lower and upper limits established for the homologous series to which the peak is assigned. See Table 9.

12.1.3. Signal-To-Noise Ratio

All ion current intensities must be >2.5 times noise level for positive identification of the PCDD/PCDF compound or a group of coeluting isomers. Figure 4 describes the procedure to be followed for the determination of the S/N.

12.1.4. Polychlorinated Diphenyl Ether Interferences

In addition to the above criteria, the identification of a GC peak as a PCDF can only be made if no signal having a S/N > 2.5 is detected, at the same retention time ( $\pm 2$  seconds), in the corresponding polychlorinated diphenyl ether (PCDPE, Table 6) channel.

12.2. For gas chromatographic peaks that have met the criteria outlined above, calculate the concentration of the PCDD or PCDF compounds using the formula:

$$C_{x} = \frac{A_{x} \times Q_{IDA}}{A_{IDA} \times W \times RRF(n)}$$

Where:

 $C_x$  = concentration of unlabeled PCDD/PCDF congeners (or group of coeluting isomers within an homologous series) usually in pg/g or pg/L,

Ax = sum of the integrated ion abundances of the quantitation ions (Table 6) for the unlabeled PCDD/PCDFs,

 $A_{IDA}$  = sum of the integrated ion abundances of the quantitation ions (Table 6) for the labeled isotope dilution analytes,

 $Q_{IDA}$  = quantity, in pg, of the isotope dilution analyte added to the sample before extraction,

W = sample size in g (if solid) or L (if liquid).

RRF(n) = Calculated mean relative response factor for the analyte

[RRF(n) with n = 1 to 17; Section 10.3.5].

If the analyte is identified as one of the 2,3,7,8-substituted PCDDs or PCDFs, RRF(n) is the value calculated using the equation in Section 10.3.5.1. However, if it is a non-2,3,7,8-substituted congener, the RRF(k) value is the one calculated using the equation in Section 10.3.5.3.2 [RRF(k) with k = 27 to 30].

12.3. Calculate the percent recovery of the nine isotope dilution analytes measured in the sample extract, using the formula:

Isotope Dilution Analytes Percent Recovery = 
$$\frac{A_{IDA} \times Q_{IS}}{Q_{IDA} \times A_{IS} \times RRF(m)} \times 100$$

Where:

 $A_{IDA}$  = sum of the integrated ion abundances of the quantitation ions (Table 6) for the labeled isotope dilution analytes,

 $A_{IS}$  = sum of the integrated ion abundances of the quantitation ions (Table 6) for the labeled internal standard; the selection of the internal standard depends on the type of congeners (see Table 5, footnotes),

 $Q_{IDA}$ = Quantity, in pg, of the isotope dilution analyte added to the sample before extraction,

 $Q_{IS}$  = Quantity, in pg, of the internal standard added to the cleaned-up sample residue before HRGC/HRMS analysis, and

RRF(m) = calculated mean relative response factor for the labeled isotope dilution analyte relative to the appropriate (see Table 5, footnotes) internal standard. This represents the mean obtained in Section 10.3.5.4 [RRF(m) with m = 18 to 26].

- 12.4. If the concentration in the final extract of any of the fifteen 2,3,7,8-substituted PCDD/PCDF compounds (Table 3) exceeds the upper method calibration limit (MCL) for that compound listed in Table 1, the linear range of response versus concentration may have been exceeded. In such cases, the following corrective actions will be undertaken:
  - 12.4.1. If the signal for the analyte has saturated the detector, a single dilution and reanalysis of the extract will be made in an attempt to bring the signal within the range of the detector. If the measured concentration of the analyte is still above the MCL, the reported concentration for the analyte will be qualified appropriately. Some programs, such as DOD QSM, require all compounds to be within the linear calibration range in which a serial dilution must be performed to achieve acceptable quantitation.

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12.4.2. If the signal for the analyte is above the MCL but does not saturate the detector, the concentration will be reported and qualified appropriately. Some programs, such as DOD QSM, require all compounds to be within the linear calibration range in which a serial dilution must be performed to achieve acceptable quantitation.

- 12.5. In either case, **with the approval of the client**, the sample may be re-extracted and/or re-analyzed with one or more of the following adjustments made to the analytical procedure in order to provide a concentration which meets client-specific data quality objectives.
  - 12.5.1. Extraction and analysis of a one tenth aliquot. This is appropriate if it will provide analyte concentration within the MCL and a representative sample aliquot.
  - 12.5.2. Extraction of an aliquot large enough to be representative with an increased concentration of isotope dilution analyte and surrogate spike components added prior to the extraction. The extract is then diluted either prior to or after the cleanup procedures.
  - 12.5.3. Dilution of the original extract. Isotope dilution analyte components are respiked at an appropriate level prior to analysis. In this case, the isotope dilution analyte recoveries are taken from the original analysis.
- 12.6. For the other congeners (including OCDD and OCDF), however, report the measured concentration and indicate that the value exceeds the upper calibration standard.
- 12.7. The total concentration for each homologous series of PCDD and PCDF is calculated by summing up the concentrations of all positively identified isomers of each homologous series. Therefore, the total should also include the 2,3,7,8-substituted congeners. The total number of GC signals included in the homologous total concentration value may be specified in the report.
- 12.8. Sample-Specific Estimated Detection Limit
  - The sample-specific estimated detection limit (EDL) or estimated quantiation limit (EQL, 8290A) is the concentration of a given analyte required to produce a signal with a peak height of at least 2.5 times the background signal level. An EDL/EQL is calculated for each 2,3,7,8-substituted congener that is not identified, regardless of whether or not other non-2,3,7,8-substituted isomers are present. Two methods of calculation can be used, as follows, depending on the type of response produced during the analysis of a particular sample.
  - 12.8.1. Samples giving a response for both quantitation ions (Tables 6 and 9) that is less than 2.5 times the background level.

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Use the expression for EDL/EQL (specific 2,3,7,8-substituted PCDD/PCDF) below to calculate an EDL/EQL for each absent 2,3,7,8-substituted PCDD/PCDF (i.e., S/N <2.5). The background level is determined by measuring the range of the noise (peak to peak) for the two quantitation ions (Table 6) of a particular 2,3,7,8-substituted isomer within an homologous series, in the region of the SICP trace corresponding to the elution of the isotope dilution analyte (if the congener possesses an isotope dilution analyte) or in the region of the SICP where the congener is expected to elute by comparison with the routine calibration data (for those congeners that do not have a <sup>13</sup>C-labeled standard), multiplying that noise height by 2.5, and relating the product to an estimated concentration that would produce that product height.

NOTE: The quantitation ions for both the unlabeled PCDDs/PCDFs and their isotope dilution analyte must be consistently paired (using either both lighter mass ions or both heavier mass ions).

Use the formula:

$$EDL_{Specific 2,3,7,8-subst.PCDD/PCDF} = \frac{2.5 \times H_x \times Q_{IDA}}{H_{IDA} \times W \times RRF(n)}$$

Where:

EDL = estimated detection limit for homologous 2,3,7,8-substituted PCDDs/PCDFs. (also EQL for Method 8290A)

 $H_x$  = height of the average noise for one of the quantitation ions (Table 6) for the unlabeled PCDDs/PCDFs.

 $H_{IDA}$  = height of one of the quantitation ions (Table 6) for the labeled isotope dilution analytes.

W, RRF (n), and  $Q_{\text{IDA}}$  retain the same meanings as defined in Section  $12.2\,$ 

12.8.2. Samples characterized by a response above the background level with a S/N of at least 2.5 for at least one of the quantitation ions (Tables 6 and 9).

When the response of a signal having the same retention times as a 2,3,7,8-substituted congener has a S/N in excess of 2.5 and does not meet any of the other qualitative identification criteria listed in Section 12.1, calculate the "Estimated Maximum Possible Concentration" (EMPC) according to the expression shown in Section 12.1, except that Ax in Section 12.1 should represent the sum of the area under the smaller peak and of the other peak area calculated using the theoretical chlorine isotope ratio. Alternatively, an EDLEQL can be calculated using the above formula and the height of one of the ions as appropriate.

12.9. The relative percent difference (RPD) is calculated as follows:

$$RPD = \frac{|S_1 - S_2|}{(S_1 + S_2)/2} \times 100$$

 $S_1$  and  $S_2$  represent sample and duplicate sample results.

- 12.10. The 2,3,7,8-TCDD toxic equivalents (TEQ) of PCDDs and PCDFs present in the sample are calculated at the data user's request. This method assigns a 2,3,7,8-TCDD toxicity equivalency factor (TEF) to each of the seventeen 2,3,7,8-substituted PCDDs and PCDFs (Table 10). The 2,3,7,8-TCDD equivalent of the PCDDs and PCDFs present in the sample is calculated by summing the TEF times their concentration for each of the compounds or groups of compounds listed in Table 10.
- 12.11. Two-GC Column TEF Determination
  - 12.11.1. The concentration of 2,3,7,8-TCDD (see note below), is calculated from the analysis of the sample extract on the 60m DB-5 fused silica capillary column. The chromatographic separation of this isomer must be  $\leq 25\%$  valley.
  - 12.11.2. For samples that have a positive result for 2,3,7,8-TCDF on the DB-5 column, the extract is reanalyzed on a 30m DB-225 fused silica column. The GC/MS conditions are altered so that only the first descriptor (Table 6) is used. The reported concentration for 2,3,7,8-TCDF is then the result above the lower calibration limit is calculated from the DB-225 analysis. The chromatographic separation between 2,3,7,8-TCDF and any other unlabeled TCDF isomers must be < 25% valley using the column performance check solution for the DB-225 column. Concentration calculations are performed as in Section 12.1 through 12.6.
  - 12.11.3. A DB-225 column can be used in the quantitative analysis of 2,3,7,8-TCDF and 2,3,7,8-TCDD analytes. Since the DB-225 cannot resolve 2,3,7,8-TCDD any positively identified 2,3,7,8-TCDD which exceeds the reporting limit shall be confirmed on a DB-5 column.
  - 12.11.4. For a gas chromatographic peak to be identified as a 2,3,7,8-substituted PCDD/PCDF congener, it must meet the ion abundance (Section 11.5.4) and signal-to-noise ratio criteria. In addition, the retention time identification criterion described in Section 11.5.4 applies here for congeners for which a carbon-labeled analog is available in the sample extract. However, the relative retention time (RRT) of the 2,3,7,8-substituted congeners for which no carbon-labeled analogs are available must fall within 0.006 units of the carbon-labeled standard RRT. Experimentally, this is accomplished by using the attributions described in Table 11 and the results from the routine

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calibration run on the DB-5 column.

#### 13. METHOD PERFORMANCE

13.1. The group/team leader has the responsibility to ensure that this procedure is performed by an associate who has been properly trained in its use and has the required expertise.

#### 13.2. Method Detection Limit

The laboratory must generate a valid method detection limit for each analyte of interest. The MDL must be below the reporting limit for each analyte. The procedure for determination of the method detection limit is given in 40 CFR Part 136, Appendix B, and further defined in SOP WS-QA-0006. MDLs are available in the Quality Assurance Department.

#### 13.3. Initial Demonstration

The laboratory must make an initial demonstration of capability for each individual method. Demonstration of capability for both soil and water matrices is required. This requires analysis of QC check samples containing all of the standard analytes for the method. For some tests it may be necessary to use more than one QC check mix to cover all analytes of interest.

- 13.3.1. Four aliquots of the QC check sample are analyzed using the same procedures used to analyze samples, including sample preparation. The concentration of the QC check sample should be less than or equivalent to the LCS samples.
- 13.3.2. Calculate the average recovery and standard deviation of the recovery for each analyte of interest. Compare these to the laboratory generated QC Limits.
- 13.4. If any analyte does not meet the acceptance criteria the test must be repeated. Only those analytes that did not meet criteria in the first test need to be evaluated. Repeated failure for any analyte indicates the need for the laboratory to evaluate the analytical procedure and take corrective action.

It must be documented that all applicable system performance criteria specified were met before analysis of any sample is performed. Table 7 provides recommended GC conditions that can be used to satisfy the required criteria. A GC column performance check is only required at the beginning of each 12-hour period during which samples are analyzed.

#### 13.5. GC Column Performance

- 13.5.1. Inject 1 or 2  $\mu$ L of the column performance check solution and acquire selected ion monitoring (SIM) data as described in Section 6.1.3 within a total cycle time of < 1 second.
- 13.5.2. The chromatographic separation between 2,3,7,8-TCDD and the peaks representing any other TCDD isomers must be resolved with a valley of  $\leq$  25

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percent (Figure 2), Where:

Valley Percent = 
$$(\frac{x}{y}) \times 100$$

x = measured as in Figure 2 from the 2,3,7,8-closest TCDD eluting isomer,

y =the peak height of 2,3,7,8-TCDD

- 13.5.3. It is the responsibility of the laboratory to verify the conditions suitable for the appropriate resolution of 2,3,7,8-TCDD from all other TCDD isomers. The GC column performance check solution also contains the known first and last PCDD/PCDF eluters under the conditions specified in this protocol. Their retention times are used for qualitative and quantitative purposes. The peak for 2,3,7,8-TCDD must be labeled on the chromatograms. The chromatograms showing the first and last eluters of a homologous series must be included.
- 13.5.4. The retention times for the switching of SIM ions characteristic of one homologous series to the next higher homologous series must be indicated in the SICP. Accurate switching at the appropriate times is absolutely necessary for accurate monitoring of these compounds.

#### 14. POLLUTION CONTROL

It is TestAmerica's policy to evaluate each method and look for opportunities to minimize waste generated (i.e., examine recycling options, ordering chemicals based on quantity needed, preparation of reagents based on anticipated usage and reagent stability). Employees must abide by the policies in Section 13 of the Corporate Environmental Health and Safety Manual (CW-E-M-001) for "Waste Management and Pollution Prevention."

#### 15. WASTE MANAGEMENT

Waste management practices are conducted consistent with all applicable rules and regulations. Excess reagents, samples and method process wastes are disposed of in an accepted manner. Waste description rules and land disposal restrictions are followed. Waste disposal procedures are incorporated by reference to SOP WS-EHS-0001. The following waste streams are produced when this method is carried out.

15.1. Autovials containing assorted solvents and extracts. As the autovials are removed from the instrument after analysis, they are collected in archive boxes and retained pending additional instructions. When no longer needed, the archive boxes are moved to the waste disposal area for disposal as PCB waste.

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## 16. REFERENCES/CROSS REFERENCES

- 16.1. SW846, Test Methods for Evaluating Solid Waste, Third edition, Update III. Method 8290 Polychlorinated Dibenzodioxins (PCDDs) and Polychlorinated Dibenzofurans (PCDFs) by high-Resolution Mass Spectrometry September 1994.
- 16.2. SW846, Test Methods for Evaluating Solid Waste, Third edition, Update IV. Method 8290A Polychlorinated Dibenzodioxins (PCDDs) and Polychlorinated Dibenzofurans (PCDFs) by high-Resolution Mass Spectrometry February 2007.
- 16.3. SW846, Test Methods for Evaluating Solid Waste, Third edition, Update III. Method 0023A, Sampling Method for Polychlorinated Dibenzo-p-dioxins and Polychlorinated Dibenzofurans Emissions from Stationary Sources. December 1996.
- 16.4. Compendium Method TO-9A "Determination of Polychlorinated, Polybrominated, and Brominated, Cholorinated Dibenxo-p-dioxins and Dibenzofurans in Ambient Air", EPA compendium of Methods for the Determination of Toxic Organic Compounds in Ambient Air, second edition, January 1997.
- 16.5. Protocol for the Analysis of 2,3,7,8-TCDD by HRGC/HRMS". J. S. Stanley and T. M. Sack, EPA 600/4-86-004.
- 16.6. "Safety in Academic Chemistry Laboratories", American Chemical Society Publication, Committee on Chemical Safety (3rd Edition, 1979.)
- 16.7. "Carcinogens Working with Carcinogens". Department of Health, Education, and Welfare, Public Health Service, Center for Disease Control. National Institute for Occupational Safety and Health. Publication No. 77-206, August 1977.
- 16.8. "OSHA Safety and Health Standards, General Industry", (29 CFR 1910) Occupational Safety and Health Administration, OSHA 2206 (revised January 1976).

## 17. METHOD MODIFICATIONS

- 17.1. Modifications from EPA 8290 and EPA 8290A
  - 17.1.1. The methods specify that 2  $\mu$ L injections are used throughout the analysis. If an instrument demonstrates adequate sensitivity and chromatographic resolution, then the analyst may use 1  $\mu$ L injections for all performance checks, standards, QC samples, and samples.
  - 17.1.2. In Section 2.7 of Method 8290 and 8290A, a retention time window of 0.005 RT units is used to tentatively identify unlabeled PCDD/PCDFs for which there are no corresponding labeled isotope dilution analytes. All available labeled isotope dilution analytes are used; therefore, a retention time window

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- of -1 to +3 seconds is used to identify all compounds. See Section 7.8.4.1 of Method 8290 and 7.9 of Method 8290A.
- 17.1.3. Tetradecane instead of nonane is used as the final solvent to increase the stability of extracts and standards. Tetradecane is less volatile than nonane. Loss of analyte as a result of solvent incompatibility is monitored through recovery checks and calibration acceptance criteria.
- 17.2. Modifications from TO-9A method
  - 17.2.1. The  $^{37}$ Cl-2,3,7,8-TCDD surrogate is present at varying levels in the calibration curve (0.5-200 pg/  $\mu$ L).
  - 17.2.2. The laboratory uses 2 labeled internal standards for the quantitation of labeled isotope dilution analytes.
  - 17.2.3. The final volume is adjusted to 20 µL in tetradecane.
  - 17.2.4. Calibration and quantitation are performed in accordance to this SOP.

#### 18. ATTACHMENTS

- 18.1. Table 1 Types of Matrices
- 18.2. Table 2 Composition of the Sample Fortification and Internal Standard Solutions.
- 18.3. Table 3 The Fifteen 2,3,7,8-Substituted PCDD and PCDF Congeners
- 18.4. Table 4 Isomers of Chlorinated Dioxins and Furans
- 18.5. Table 5 Concentrations of Calibration Solutions
- 18.6. Table 6 Ions Monitored for PCDDs/PCDFs
- 18.7. Table 7 Recommended GC Operating Conditions
- 18.8. Table 8 Congeners in the GC Performance Evaluation Solution (DB-5)
- 18.9. Table 9 Theoretical Ion Abundance Ratios and Control Limits
- 18.10. Table 10 2,3,7,8-TCDD Equivalent Factors
- 18.11. Table 11 TEF: Analyte Relative Retention Time Reference Attributes
- 18.12. Figure 1 Compound Structure

- 18.13. Figure 2 GC Performance Check Chromatogram on the DB-5 Column
- 18.14. Figure 3 PFK Peak Profile
- 18.15. Figure 4 Manual Determination of Signal-to-Noise
- 18.16. Appendix A Periodic Wipe Test Performance

## 19. REVISION HISTORY

- 19.1. WS-ID-0005, Revision 7.5, Effective 04/19/2013
  - 19.1.1. Replaced all instances of 'internal standard' with isotope dilution analyte' and all instances of 'recovery standard' with 'internal standard' to conform with TALS naming guidelines.
  - 19.1.2. Editorial revisions.
- 19.2. WS-ID-0005, Revision 7.4, Effective 01/14/2011.
  - 19.2.1. Editorial revisions.
- 19.3. WS-ID-0005, Revision 7.3, Effective 12/30/2009
  - 19.3.1. Editorial revisions.
- 19.4. WS-ID-0005, Revision 7.2, Effective 11/02/2009
  - 19.4.1. Section 6.1: Inserted "Preventive and routine maintenance is described in the 'Schedule of Routine Maintenance' in the QAM."
  - 19.4.2. Section 12.1.2: Removed the word "presumptive" and inserted "above the lower calibration limit" after the word result.

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TABLE 1

Types of Matrices, Sample Sizes and 2,3,7,8-TCDD-Based
Method Calibration Limits (Parts per Trillion)

	Water	Soil Sediment Paper Pulp	Fly Ash	Human/ Fish Tissue	Adipose Tissue	Sludges, Fuel Oil	Still- Bottom	Ambient or Source Samples
Lower MCL(a)	0.01	1.0	2.0	1.0	2.0	10	20	40
Upper MCL(a)	4.0	400	400	400	400	2000	4000	8000
Weight (g)	1000	10	10	10	10	2.0	1.0	1 sample
IDA Spiking Levels (ng)	2.0	2.0	2.0	2.0	2.0	2.0	2.0	4.0
Final Extract Volume (μL)	20	20	20	20	20	20	20	20

<sup>(</sup>a) For other congeners, multiply the values by 1 for TCDF, by 5 for PeCDD/PeCDF/HxCDD/HxCDF/HpCDD/HpCDF, and by 10 for OCDD/OCDF.

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TABLE 2

Composition of the Sample Fortification and Internal Standard Solutions

Analyte	Sample Fortification Solution	Internal Standard Solution
	Concentration pg/µL;	Concentration pg/μL;
	Solvent: Isooctane	Solvent: Tetradecane
<sup>13</sup> C-2,3,7,8-TCDD	2 <sup>(a)</sup> , 100 <sup>(c)</sup> 2 <sup>(a)</sup> , 100 <sup>(c)</sup>	
<sup>13</sup> C -2,3,7,8-TCDF	2 <sup>(a)</sup> , 100 <sup>(c)</sup>	
<sup>13</sup> C -1,2,3,4-TCDD		100
<sup>13</sup> C -1,2,3,7,8-PeCDD	2 <sup>(a)</sup> , 100 <sup>(c)</sup>	
<sup>13</sup> C -1,2,3,7,8-PeCDF	2 <sup>(a)</sup> , 100 <sup>(c)</sup>	
<sup>13</sup> C -1,2,3,6,7,8-HxCDD	2 <sup>(a)</sup> , 100 <sup>(c)</sup>	
<sup>13</sup> C -1,2,3,4,7,8-HxCDF <sup>(d)</sup>	2 <sup>(a)</sup> , 100 <sup>(c)</sup>	
<sup>13</sup> C -1,2,3,7,8,9-HxCDD		100
37GL 2.2.7.0. TGDD(b)(c)	0.8 <sup>(b),</sup> 100 <sup>(c)</sup>	
<sup>37</sup> Cl-2,3,7,8-TCDD <sup>(b)(c)</sup>		
13 G 2 2 4 7 0 D GD 7(c)	100 <sup>(c)</sup>	
<sup>13</sup> C -2,3,4,7,8-PeCDF <sup>(c)</sup>	100 <sup>(c)</sup>	
<sup>13</sup> C -1,2,3,6,7,8-HxCDF <sup>(c)(d)</sup>	100 <sup>(c)</sup>	
<sup>13</sup> C -1,2,3,4,7,8-HxCDD <sup>(c)</sup>	100 <sup>(c)</sup>	
<sup>13</sup> C -1,2,3,4,7,8,9-HpCDD <sup>(c)</sup>	100 <sup>(c)</sup>	
<sup>13</sup> C -1,2,3,4,6,7,8-HpCDD	$2^{(a)}$ , $100^{(c)}$	
<sup>13</sup> C -1,2,3,4,6,7,8-HpCDF	2 <sup>(a)</sup> , 100 <sup>(c)</sup>	
<sup>13</sup> C -OCDD	$4^{(a)}, 200^{(c)}$	

- (a) Standard 8290, 8290A, Method 23, Method 0023A, TO9 and TO9A Sample Fortification Solution concentrations
- (b) Method TO9 and TO9A surrogate concentrations
- (c) Method 23 and Method 0023A surrogate concentrations
- (d) <sup>13</sup>C-1,2,3,6,7,8-HxCDF is used as a Sample Fortification Solution and <sup>13</sup>C -1,2,3,4,7,8-HxCDF is used as a surrogate solution in Method 0023A

TABLE 3

The Seventeen 2,3,7,8-Substituted PCDD and PCDF Congeners

PCDD	PCDF
2,3,7,8-TCDD(*)	2,3,7,8-TCDF(*)
1,2,3,7,8-PeCDD(*)	1,2,3,7,8-PeCDD(*)
1,2,3,6,7,8-HxCDD(*)	2,3,4,7,8-PeCDF
1,2,3,4,7,8-HxCDD	1,2,3,6,7,8-HxCDF
1,2,3,7,8,9-HxCDD(+)	1,2,3,7,8,9-HxCDF
1,2,3,4,6,7,8-HpCDD(*)	1,2,3,4,7,8-HxCDF(*)
1,2,3,4,5,6,7,8-OCDD(*)	2,3,4,6,7,8-HxCDF
	1,2,3,4,6,7,8-HpCDF(*)
	1,2,3,4,7,8,9-HpCDF
	1,2,3,4,5,6,7,8-OCDF

<sup>(\*)</sup>The <sup>13</sup>C -labeled analog is used as an isotope dilution analyte. (+)The <sup>13</sup>C -labeled analog is used as a internal standard.

TABLE 4

Isomers of Chlorinated Dioxins and Furans as a Function of the Number of Chlorine Atoms

# of Chlorine Atoms	# of Dioxin Isomers	# of 2,3,7,8 Isomers	# of Furan Isomers	# of 2,3,7,8 Isomers
1	2		4	
2	10		16	
3	14		28	
4	22	1	38	1
5	14	1	28	2
6	10	3	16	4
7	2	1	4	2
8	1	1	1	1
Total	75	7	135	10

TABLE 5
High Resolution Concentration Calibration Solutions

	Compound		Con	centration (n	g/mL)	
RRF		CS2	CS3	CS4	CS5	CS6
(n)(m)				(ICV(6))		
	Native CDDs and CDFs					
1	2,3,7,8-TCDD	0.5	2	10	40	200
2	2,3,7,8-TCDF	0.5	2	10	40	200
3	1,2,3,7,8-PeCDD	2.5	10	50	200	1000
4	1,2,3,7,8-PeCDF	2.5	10	50	200	1000
5	2,3,4,7,8-PeCDF	2.5	10	50	200	1000
6	1,2,3,4,7,8-HxCDD	2.5	10	50	200	1000
7	1,2,3,6,7,8-HxCDD	2.5	10	50	200	1000
8	1,2,3,7,8,9-HxCDD	2.5	10	50	200	1000
9	1,2,3,4,7,8-HxCDF	2.5	10	50	200	1000
10	1,2,3,6,7,8-HxCDF	2.5	10	50	200	1000
11	1,2,3,7,8,9-HxCDF	2.5	10	50	200	1000
12	2,3,4,6,7,8-HxCDF	2.5	10	50	200	1000
13	1,2,3,4,6,7,8-HpCDD	2.5	10	50	200	1000
14	1,2,3,4,6,7,8-HpCDF	2.5	10	50	200	1000
15	1,2,3,4,7,8,9-HpCDF	2.5	10	50	200	1000
16	OCDD	5.0	20	100	400	2000
17	OCDF	5.0	20	100	400	2000
	Labeled CDDs and CDFs					
18	<sup>13</sup> C <sub>12</sub> -2,3,7,8-TCDD	100	100	100	100	100
19	<sup>13</sup> C <sub>12</sub> -2,3,7,8-TCDF	100	100	100	100	100
20	<sup>13</sup> C <sub>12</sub> -1,2,3,7,8-PeCDD	100	100	100	100	100
21	<sup>13</sup> C <sub>12</sub> -1,2,3,7,8-PeCDF	100	100	100	100	100
	<sup>13</sup> C <sub>12</sub> -2,3,4,7,8-PeCDF	100	100	100	100	100
	<sup>13</sup> C <sub>12</sub> -1,2,3,4,7,8-HxCDD	100	100	100	100	100
22	<sup>13</sup> C <sub>12</sub> -1,2,3,6,7,8-HxCDD	100	100	100	100	100
23	<sup>13</sup> C <sub>12</sub> -1,2,3,4,7,8-HxCDF	100	100	100	100	100
	<sup>13</sup> C <sub>12</sub> -1,2,3,6,7,8-HxCDF	100	100	100	100	100
	<sup>13</sup> C <sub>12</sub> -1,2,3,7,8,9-HxCDF	100	100	100	100	100
	<sup>13</sup> C <sub>12</sub> 2,3,4,6,7,8-HxCDF	100	100	100	100	100
24	$^{13}C_{12}$ -1,2,3,4,6,7,8-	100	100	100	100	100
	HpCDD					
25	$^{13}$ C <sub>12</sub> -1,2,3,4,6,7,8-	100	100	100	100	100
	HpCDF					
	<sup>13</sup> C <sub>12</sub> -1,2,3,4,7,8,9-	100	100	100	100	100

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	Compound	Concentration (ng/mL)				
RRF (n)(m)		CS2	CS3	CS4 (ICV(6))	CS5	CS6
	HpCDF					
26	<sup>13</sup> C <sub>12</sub> -OCDD	200	200	200	200	200
	Cleanup Standard/ FS					
	<sup>37</sup> Cl <sub>4</sub> 2,3,7,8-TCDD	0.5	2	10	40	200
	Internal Standards					
	<sup>13</sup> C <sub>12</sub> -1,2,3,4-TCDD	100	100	100	100	100
	<sup>13</sup> C <sub>12</sub> -1,2,3,7,8,9-HxCDD	100	100	100	100	100

TABLE 6\*
Elemental Compositions and Exact Masses of the Ions
Monitored by HR/MS for PCDD's and PCDF's

Descriptor	Exact m/z (1)	m/z Type	Elemental Composition	Substance (2)
1	292.9825	QC	$C_7F_{11}$	PFK
	303.9016	M	C <sub>12</sub> H <sub>4</sub> <sup>35</sup> Cl <sub>4</sub> O	TCDF
	305.8987	M+2	$C_{12}H_4^{35}Cl_3^{37}ClO$	TCDF
	315.9419	M	$^{13}\text{C}_{12}\text{H}_4$ $^{35}\text{Cl}_4\text{O}$	TCDF (3)
	317.9389	M+2	<sup>13</sup> C <sub>12</sub> H <sub>4</sub> <sup>35</sup> Cl <sub>3</sub> <sup>37</sup> ClO	TCDF (3)
	319.8965	M	$C_{12}H_4^{35}Cl_4O_2$	TCDD
	321.8936	M+2	$C_{12}H_4^{35}Cl_3^{37}ClO_2$	TCDD
	327.8847	M	$C_{12}H_4^{37}Cl_4O_2$	TCDD (4)
	330.9792	Lock	$C_7F_{13}$	PFK
	331.9368	M	$^{13}\text{C}_{12}\text{H}_4{}^{35}\text{Cl}_4\text{O}_2$	TCDD (3)
	333.9339	M+2	$^{13}\text{C}_{12}\text{H}_4^{\ 35}\text{Cl}_3^{\ 37}\text{ClO}_2$	TCDD (3)
	339.8597	M+2	$C_{12}H_3^{35}Cl_4^{37}ClO$	PeCDF
	341.8567	M+4	$C_{12}H_3^{35}Cl_3^{37}ClO$	PeCDF
	375.8364	M+2	$C_{12}H_4^{35}Cl_5^{37}ClO$	HxCDPE
	409.7974	M+2	$C_{12}H_3$ $^{35}Cl_6$ $^{37}ClO$	HpCDPE
2	330.9792	QC	$C_{7}F_{13}$	PFK
	339.8597	M+2	$C_{12}H_3^{35}Cl_4^{37}ClO$	PeCDF
	341.8567	M+4	$C_{12}H_3^{35}Cl_3^{37}Cl_2O$	PeCDF
	342.9792	Lock	$C_8F_{12}$	PFK
	351.9000	M+2	$^{13}\text{C}_{12}\text{H}_3{}^{35}\text{Cl}_4{}^{37}\text{ClO}$	PeCDF
	353.8970	M+4	$^{13}\text{C}_{12}\text{H}_3^{\ 35}\text{Cl}_4^{\ 37}\text{ClO}$	PeCDF (3)
	354.9792	Lock	$C_9F_{13}$	PFK
	355.8546	M+2	$C_{12}H_3^{35}Cl_4^{37}ClO_2$	PeCDD
	357.8516	M+4	$C_{12}H_3^{35}Cl_3^{37}Cl_2O_2$	PeCDD
	366.9793	QC	$C_9F_{13}$	PFK
	367.8949	M+2	$^{13}\text{C}_{12}\text{H}_3^{\ 35}\text{Cl}_4^{\ 37}\text{ClO}_2$	PeCDD (3)
	369.8919	M+4	$^{13}\text{C}_{12}\text{H}_3^{\ 35}\text{Cl}_3^{\ 37}\text{Cl}_2\text{O}_2$	PeCDD (3)
	409.7974	M+2	$C_{12}H_3^{35}Cl_6^{37}ClO$	HpCDPE
3	373.8208	M+2	$C_{12}H_2^{35}Cl_5^{37}ClO$	HxCDF
	375.8178	M+4	$C_{12}H_2^{35}Cl_4^{37}Cl_2O$	HxCDF
	380.9760	Lock	$C_8F_{15}$	PFK
	383.8639	M	$^{13}\text{C}_{12}\text{H}_2^{35}\text{Cl}_6\text{O}$	HxCDF (3)
	385.8610	M+2	$^{13}\text{C}_{12}\text{H}_2^{\ 35}\text{Cl}_5^{\ 37}\text{ClO}$	HxCDF (3)
	389.8157	M+2	$C_{12}H_2^{35}Cl_5^{37}ClO_2$	HxCDD
	391.8127	M+4	$C_{12}H_2^{35}Cl_4^{37}Cl_2O_2$	HxCDD
	392.9760	Lock	C <sub>9</sub> F <sub>15</sub>	PFK
	401.8559	M+2	$^{13}\text{C}_{12}\text{H}_2^{\ 35}\text{Cl}_5^{\ 37}\text{ClO}_2$	HxCDD (3)
	403.8529	M+4	$^{13}\text{C}_{12}\text{H}_2^{\ 35}\text{Cl}_4^{\ 37}\text{Cl}_2\text{O}_2$	HxCDD (3)
ļ	430.9728	QC	C <sub>0</sub> F <sub>17</sub>	PFK
ļ	445.7550	M+4	$C_{12}H_2^{35}Cl_6^{37}Cl_2O$	OCDPE
4	392.9760	QC	$C_9F_{15}$	PFK
	407.7818	M+2	$C_{12}H^{35}Cl_6^{37}ClO$	HpCDF
ļ	409.7789	M+4	$C_{12}H^{35}Cl_5^{37}Cl_2O$	HpCDF

Descriptor	Exact m/z (1)	m/z Type	Elemental Composition	Substance (2)
	417.8253	M	$^{13}\text{C}_{12}\text{H}^{35}\text{Cl}_7\text{O}$	HpCDF (3)
	419.8220	M+2	$^{13}\text{C}_{12}\text{H}^{35}\text{Cl}_6^{\ 37}\text{ClO}$	HpCDF (3)
	423.7766	M+2	$C_{12}H^{35}Cl_6^{37}ClO_2$	HpCDD
	425.7737	M+4	$C_{12}H^{35}Cl_5^{37}Cl_2O_2$	HpCDD
	430.9729	Lock	C <sub>9</sub> F <sub>17</sub>	PFK
	435.8169	M+2	$^{13}\text{C}_{12}\text{H}^{35}\text{Cl}_6^{37}\text{ClO}_2$	HpCDD (3)
	437.8140	M+4	$^{13}\text{C}_{12}\text{H}^{35}\text{Cl}_5^{37}\text{CL}_2\text{O}_2$	HpCDD (3)
	479.7165	M+4	$C_{12}H^{35}Cl_7^{37}Cl_2O$	NCDPE
5	392.9760	QC	$C_9F_{15}$	PFK
	441.7428	M+2	$C_{12}^{35}Cl_7^{37}ClO$	OCDF
	442.9728	Lock	$C_{10}F_{17}$	PFK
	443.7399	M+4	$C_{12}^{35}Cl_6^{37}Cl_2O$	OCDF
	457.7377	M+2	$C_{12}^{35}Cl_7^{37}ClO_2$	OCDD
	459.7348	M+4	$C_{12}^{35}Cl_6^{37}Cl_2O_2$	OCDD
	469.7779	M+2	$^{13}\text{C}_{12}^{35}\text{Cl}_7^{37}\text{ClO}_2$	OCDD (3)
	471.7750	M+4	$^{13}\text{C}_{12}^{35}\text{Cl}_{6}^{37}\text{Cl}_{2}\text{O}_{2}$	OCDD (3)
	479.7165	M+4	$C_{12}Cl_8^{37}Cl_2O$	NCDPE
	513.6775	M+4	$^{13}\text{C}_{12}^{35}\text{Cl}_{8}^{37}\text{Cl}_{2}\text{O}$	DCDPE

(a) The following nuclidic masses were used:

H = 1.007825 O = 15.994915 C = 12.000000 O = 15.994915  = 15.

F = 18.9984

S = Isotope dilution analyte/internal standard

<sup>\*</sup>The homologous groups for functions 1-3 do not use the same lockmass as described in Table 6. They use masses 316.9824, 366.9792, and 380.9760, respectively.

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#### TABLE 7

# **Recommended GC Operating Conditions**

The GC Operating Conditions (Temperatures (°C), and Times (minutes)) Are as Follows:

Injector Temperature: 280°C Interface Temperature: 280°C

Initial Temperature and Time: 190°C / 1 Minute

Temperature Program: 190°C, increasing at a rate of 4°C per minute up to 240°C, and maintaining at this temperature until the last tetra of the tetra- group has eluted from the column. (The total time required for this is approximately 25 minutes, depending on the length of the column). The maintained temperature of 240°C is then increased to 320°C at the rate of 20°C per minute and held at this level until the last compound (octa-group) has eluted from the column.

TABLE 8

PCDD and PCDF Congeners Present in the GC Performance Evaluation Solution and Used for Defining the Homologous GC Retention Time Windows on a 60-M DB-5 Column (b)

# of Chlorine	PCDD Positional Isomer		PCDF Positional Isomer	
Atoms	Early Eluter	Late Eluter	Early Eluter	Late Eluter
4 <sup>(a)</sup>	1,3,6,8	1,2,8,9	1,3,6,8	1,2,8,9
5	1,2,4,6,8/1,2,4,7,9	1,2,3,8,9	1,3,4,6,8	1,2,3,8,9
6	1,2,3,4,6,8	1,2,3,4,6,7	1,2,3,4,6,8	1,2,3,4,8,9
7	1,2,3,4,6,7,8	1,2,3,4,6,7,9	1,2,3,4,6,7,8	1,2,3,4,6,7,9
8	1,2,3,4,6,7,8,9		1,2,3,4	,6,7,8,9

<sup>(</sup>a) In addition to these two PCDD isomers, the 1,2,3,4-, 1,2,3,7-, 1,2,3,8-, 2,3,7,8-, <sup>13</sup>C<sub>12</sub>-2,3,7,8-, and 1,2,3,9-TCDD isomers must also be present.

- (b) The PCDF Congeners present in GC the Performance Evaluation Solution for the 30 m DB-225 column include:
  - 1,2,3,9-TCDF
  - 2,3,7,8-TCDF
  - 2,3,4,7-TCDF
  - ${}^{13}C_{12}$ -2,3,7,8-TCDF

Column performance criteria is met when the percent valleys between the 2,3,7,8-TCDF analyte and the closest eluting isomers are < 25%.

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TABLE 9

Theoretical Ion Abundance Ratios and Their Control Limits for PCDDs and PCDFs

# of Chlorine	Ion Type	Theoretical Ratio	Contro	Limits
Atoms			Lower	Upper
4	M / M+2	0.77	0.65	0.89
5	M+2 / M+4	1.55	1.32	1.78
6	M+2 / M+4	1.24	1.05	1.43
$6^{(a)}$	M / M+2	0.51	0.43	0.59
7 <sup>(b)</sup>	M/M+2	0.44	0.37	0.51
7	M+2 / M+4	1.04	0.88	1.20
8	M+2 / M+4	0.89	0.76	1.02

(a) Used only for <sup>13</sup>C-HxCDF (IS)

(b) Used only for <sup>13</sup>C-HpCDF (IS)

TABLE 10

2,3,7,8-TCDD Equivalent Factors (TEFs) for the Polychlorinated
Dibenzodioxins and Dibenzofurans

Number	Compound(s)	TEF
1	2,3,7,8-TCDD	1.00
2	1,2,3,7,8-PeCDD	0.50
3	1,2,3,6,7,8-HxCDD	0.10
4	1,2,3,7,8,9-HxCDD	0.10
5	1,2,3,4,7,8-HxCDD	0.10
6	1,2,3,4,6,7,8-HpCDD	0.01
7	OCDD	0.001
8	2,3,6,7-TCDF	0.1
9	1,2,3,7,8-PeCDF	0.05
10	2,3,4,7,8PeCDF	0.5
11	1,2,3,6,7,8-HxCDF	0.1
12	1,2,3,7,8,9-HxCDF	0.1
13	1,2,3,4,7,8-HxCDF	0.1
14	2,3,4,6,7,8-HxCDF	0.1
15	1,2,3,4,6,7,8-HpCDF	0.01
16	1,2,3,4,7,8,9-HpCDF	0.01
17	OCDF	0.001

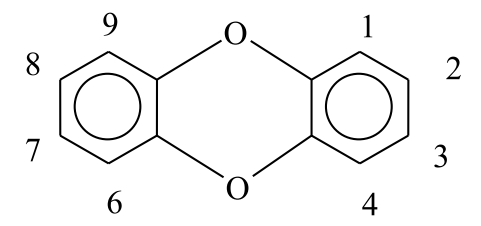
**TABLE 11** 

# Toxicity Equivalency Factor: Analyte Relative Retention Time Reference Attributes

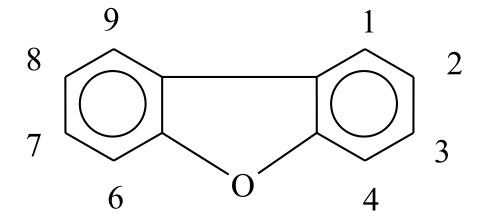
Analyte	Analyte RRT Reference (a)
1,2,3,4,7,8-HxCDD	<sup>13</sup> C <sub>12-</sub> 1,2,3,6,7,8-HxCDD
1,2,3,6,7,8-HxCDF	<sup>13</sup> C <sub>12</sub> .1,2,3,4,7,8-HxCDF
1,2,3,7,8,9-HxCDF	<sup>13</sup> C <sub>12</sub> .1,2,3,4,7,8-HxCDF
2,3,4,6,7,8-HxCDF	<sup>13</sup> C <sub>12-</sub> 1,2,3,4,7,8-HxCDF

<sup>(</sup>a) The retention time of 2,3,4,7,8-PeCDF on the DB-5 column is measured relative to <sup>13</sup>C<sub>12-1</sub>,3,7,8-PeCDF and the retention time of 1,2,3,4,7,8,9-HpCDF relative to <sup>13</sup>C<sub>12-1</sub>,2,3,4,6,7,8-HpCDF

**FIGURE 1** Structure of Dibenzodioxin and Dibenzofuran



Dibenzodioxin



Dibenzofuran

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FIGURE 2

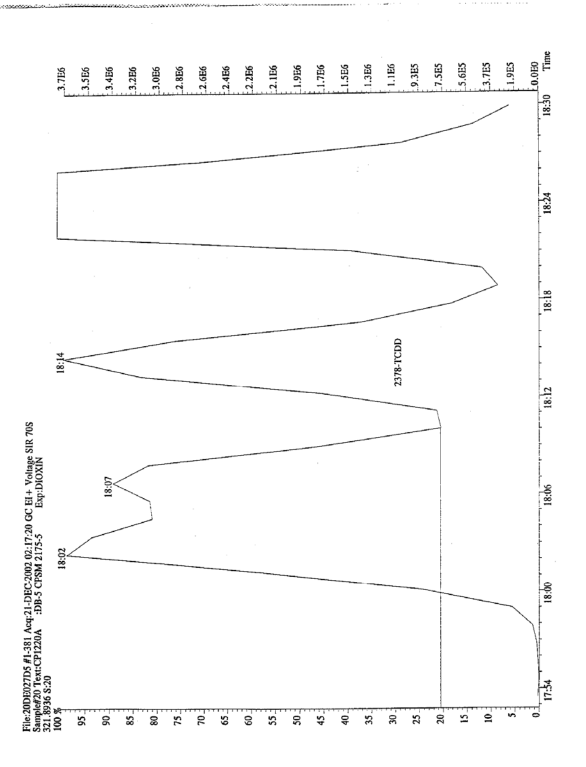
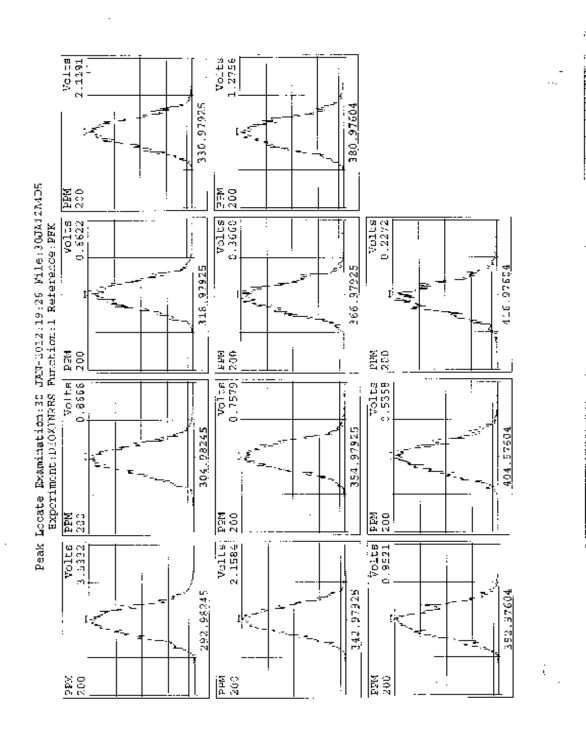
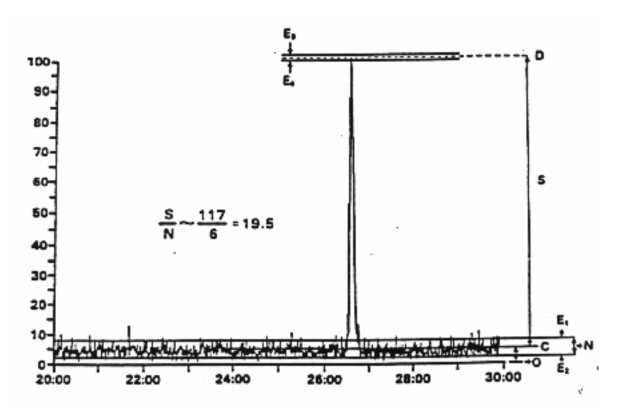


Figure 3



# FIGURE 4



# Manual determination of S/N.

The peak height (S) is measured between the mean noise (lines C and D). These mean signal values are obtained by tracing the line between the baseline average noise extremes, El and E2, and between the apex average noise extremes. E3 and E4, at the apex of the signal.

<u>HOTE</u>: It is imperative that the instrument interface amplifier electronic zero offset be set high enough so that negative going baseline noise is recorded.

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#### APPENDIX A

This procedure is designed for the periodic evaluation of potential contamination by 2,3,7,8-substituted PCDD/PCDF congeners of the working areas inside the laboratory.

#### PERFORMING WIPE TEST

Perform the wipe tests on surface areas of two inches by one foot with laboratory wipers saturated with distilled-in-glass acetone or appropriate solvent using a pair of clean stainless steel forceps. Use one wiper for each of the designated areas. Combine the wipers to one composite sample in an extraction jar containing 200 mL distilled-in-glass hexane. Place an equal number of unused wipers in 200 mL hexane and use this as a control

#### SAMPLE PREPARATION

Close the jar containing the wipes and 200 mL hexane and extract for 20 minutes using a wrist-action shaker. Use an appropriate means to reduce the volume to approximately 1.0 mL. Put through an alumina column to clean up potential interfering compounds. Add appropriate amount of internal standard.

## **EXTRACT ANALYSIS**

Concentrate the contents of the vial to a final volume of 20  $\mu$ L (either in a minivial or in a capillary tube). Inject 2  $\mu$ L of each extract (wipe and control) onto a capillary column and analyze for 2,3,7,8-substituted PCDDs/PCDFs as specified in the analytical method Section 11 (this exhibit). Perform calculations according to Section 12 (this exhibit).

## REPORTING FORMAT

Report the presence of 2,3,7,8-substituted PCDDs and PCDFs as a quantity (pg or ng) per wipe test experiment (WTE). Under the conditions outlined in this analytical protocol, a lower limit of calibration of 25 pg/WTE is expected for 2,3,7,8-TCDD. A positive response for the blank (control) is defined as a signal in the TCDD retention time window at any of the masses monitored which is equivalent to or above 8 pg of 2,3,7,8-TCDD per WTE. For other congeners, use the multiplication factors listed in Table 1, footnote (a) (e.g., for OCDD, the lower MCL is  $25 \times 5 = 125 \text{ pg/WTE}$  and the positive response for the blank would be  $8 \times 5 = 40 \text{ pg}$ ). Also, report the recoveries of the isotope dilution analytes during the simplified cleanup procedure.

## FREQUENCY OF WIPE TESTS

Wipe tests should be performed when there is evidence of contamination in the method blanks.

CORRECTIVE ACTION

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An upper limit of 25 pg per TCDD isomer and per wipe test experiment is allowed. (Use multiplication factors listed in footnote (a) from Table 1 for other congeners.) This value corresponds to the lower calibration limit of the analytical method. Steps to correct the contamination must be taken whenever these levels are exceeded. To that effect, first vacuum the working places (hoods, benches, sink) using a vacuum cleaner equipped with a high-efficiency particulate absorbent (HEPA) filter and then wash with a detergent. A new set of wipes should be analyzed before anyone is allowed to work in the dioxin area of the laboratory.

The test results and the decontamination procedure must be reviewed with EH&S.

## **West Sacramento**



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`Title:

Preparation of Samples for Analysis of Polychlorinated Dioxins and Furans for Analysis HRGC/HRMS

[Methods 8290, 8290A & TO-9A]

	Approvals (S	Signature/Date):
Elizabeth Ngu/en Technical Manager	/2/2///2 /Date	Joe/Schairer Date Health & Safety Manager / Coordinator
Maril Afflus you Williams Weir Quality Assurance Manager	12/20/12   Date	Marka Buechler Date Laboratory Director

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## 1. SCOPE AND APPLICATION

- 1.1. This method provides procedures for the preparation of samples prior to the analysis of 2,3,7,8-tetrachlorodibenzo-p-dioxin (2,3,7,8-TCDD), polychlorinated dibenzo-p-dioxins (tetra- through octachlorinated homologs; PCDDs), and polychlorinated dibenzofurans (tetra- through octachlorinated homologs; PCDFs) in a variety of environmental matrices at part-per-trillion (ppt) concentrations by SW 846 Method 8290. The analytical method calls for the use of high-resolution gas chromatography and high-resolution mass spectrometry (HRGC/HRMS) on purified sample extracts. Refer to Table 1 for the list of analytes. Analysis is by SOP WS-ID-0005.
- 1.2. The sensitivity of this method is dependent upon the level of interferences within a given matrix.
- 1.3. This method is designed for use by analysts who are experienced with residue analysis.
- 1.4. Samples containing concentrations of specific congeners (PCDDs and PCDFs) that are greater than the calibration limit should be analyzed by a protocol designed for such concentrations, such as 8280A/B.

## 2. SUMMARY OF METHOD

- 2.1. This procedure uses matrix-specific extraction and analyte-specific cleanup techniques.
- 2.2. A specified amount (see Table 1) of soil, sediment, fly ash, water, sludge (including paper pulp), still-bottom, fuel oil, chemical reactor residue, air sample (QFF, PUF or XAD media) or fish tissue, is spiked with a solution containing specified amounts of each of nine isotopically (<sup>13</sup>C) labeled PCDDs/PCDFs listed in Table 2. The sample is then extracted according to a matrix-specified extraction procedure. The extraction procedures are: a) toluene Soxhlet (or equivalent) extraction, for soil, sediment, fly ash samples, aqueous sludges, and solid air matrices (XAD, QFF, PUF); b) methylene chloride liquid-liquid extraction or solid phase extraction for water samples; c) dilution of a small sample aliquot in solvent for wastes/chemical products; and d) toluene (or hexane/methylene chloride) Soxhlet (or equivalent) extraction for fish tissue. This method can also use solid phase extraction (SPE), however, Test America West Sacramento is in the developmental stages for this extraction type and is not currently certified for its use.
- 2.3. If interferences are present, extracts may be cleaned as described below. The extracts are submitted to an acid and/or base washing treatment and dried. Following a solvent exchange step, the residue is cleaned up by column chromatography on acid/base silica, acid alumina and carbon on silica. The preparation of the final extract for HRGC/HRMS analysis is accomplished by adding 20  $\mu$ L of a tetradecane solution containing 100 pg/ $\mu$ L of each of the two recovery standards  $^{13}C_{12}$ -1,2,3,4-TCDD and

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<sup>13</sup>C<sub>12</sub> -1,2,3,7,8,9-HxCDD (Table 2) to the concentrated eluate. The former is used to determine the percent recoveries of tetra- and penta-chlorinated PCDD/PCDF internal standards while the latter is used for the determination of hexa-, hepta- and octa-chlorinated PCDD/PCDF internal standard percent recoveries. Upon client approval, less final volume can be used to decrease detection limit and more final volume can be used to decrease severe interferences.

## 3. **DEFINITIONS**

- 3.1. Definitions of terms used in this SOP may be found in the glossary of the Quality Assurance Manual (QAM).
- 3.2. Data qualifiers are defined on each data report. Commonly used data qualifiers are defined in the QAM.
- 3.3. Internal Standard: An internal standard is a <sup>13</sup>C-labeled analog of a congener chosen from the compounds listed in Table 2. Internal standards are added to all samples including method blanks and quality control samples before extraction, and they are used to quantitate the concentration of the analytes. Nine internal standards are used in this method. There is one for each of the dioxin and furan homologs (except for OCDF) with the degree of chlorination ranging from four to eight. Additional internal standards may be added to act as retention time references, but they are not used for quantitation.
- 3.4. Recovery Standard: Two recovery standards are used to determine the percent recoveries for the internal standards. The <sup>13</sup>C<sub>12</sub>-1,2,3,4-TCDD is used to measure the percent recoveries of the tetra- and pentachlorinated internal standards while <sup>13</sup>C<sub>12</sub>-1,2,3,7,8,9-HxCDD is used to determine the recovery of the hexa-hepta- and octachlorinated internal standards. <sup>13</sup>C<sub>12</sub>-1,2,3,7,8,9-HxCDD also acts as a retention time reference for the unlabeled analog present in sample extracts. They are added to the final sample extract before HRGC/HRMS instrument analysis.
- 3.5. Cleanup Recovery Standard (CRS): A <sup>37</sup>Cl<sub>4</sub>-2,3,7,8-TCDD analog that is added to each sample following extraction to measure the efficiency of the cleanup process.

#### 4. INTERFERENCES

- 4.1. Solvents, reagents, glassware and other sample processing hardware may yield discrete artifacts or elevated baselines that may cause misinterpretation of the chromatographic data. All of these materials must be demonstrated to be free from interferents under the conditions of analysis by running laboratory method blanks. Analysts shall not use PVC gloves.
- 4.2. The use of high-purity reagents and solvents helps minimize interference problems. Purification of solvents by distillation in all-glass systems may be necessary.

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4.3. Proper cleaning of glassware is extremely important because glassware may not only contaminate the samples, but may also remove the analytes of interest by adsorption on the glassware surface.

- 4.3.1. Glassware should be rinsed with solvent and washed with a detergent solution as soon after use as is practical. Sonication of glassware containing a detergent solution for approximately 30 seconds may aid in cleaning. Glassware with removable parts, particularly separatory funnels with Teflon stopcocks, must be disassembled prior to detergent washing.
- 4.3.2. After detergent washing, glassware should be immediately rinsed with acetone, toluene, hexane, and then methylene chloride.
- 4.3.3. Do not kiln reusable glassware in an oven as a routine part of cleaning. Kilning may be warranted after particularly dirty samples are encountered, but should be minimized, as repeated kilning of glassware may cause the formation of active sites on the glass surface that will irreversibly adsorb PCDDs/ PCDFs.
- 4.3.4. Immediately prior to use, Soxhlet (or equivalent) extraction glassware should be pre-extracted with toluene for a minimum of 3 hours. Note:

  Accelerated extractors such as the Soxtherm can use a shorter cleaning cycle which exhibits subsequent extractions free of cross contamination and interferences

Note: Re-use of glassware should be minimized to avoid the risk of contamination. All glassware that is re-used must be scrupulously cleaned as soon as possible after use, applying the following procedure:

- 4.4. Interferences co-extracted from samples will vary considerably from source to source, depending on the diversity of the site being sampled. Interfering compounds may be present at concentrations several orders of magnitude higher than the PCDDs and PCDFs. The most frequently encountered interferences are chlorinated-biphenyls, methoxy biphenyls, hydroxy biphenyl ethers, benzyl phenyl ethers, polynuclear aromatics, and pesticides. Because very low levels of PCDDs and PCDFs are measured by this method, the elimination of interferences is essential. The cleanup steps given in Sections 11.12 thru 11.16 can be used to reduce or eliminate these interferences.
  - 4.4.1. If South Carolina samples show diphenyl ethers at levels that could contribute to positive furan hits, a subsequent clean-up to remove them must be performed.

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## 5. SAFETY

Employees must abide by the policies and procedures in the Corporate Environmental Health and Safety Manual (CW-E-M-001), the West Sacramento Addendum to the Corporate EH&S Manual (WS-PEHS-002) and this document. This procedure may involve hazardous material, operations and equipment. This SOP does not purport to address all of the safety problems associated with its use. It is the responsibility of the user of the method to follow appropriate safety, waste disposal and health practices under the assumption that all samples and reagents are potentially hazardous. Safety glasses, gloves, lab coats and closed-toes, nonabsorbent shoes are a minimum.

- 5.1. Specific Safety Concerns or Requirements
  - 5.1.1. Hearing protection must be worn when using mechanical systems to grind fish, tissue, or paper/pulp samples.
  - 5.1.2. Finely divided dry soils contaminated with PCDDs and PCDFs are particularly hazardous because of the potential for inhalation and ingestion. Such samples are to be processed in a confined environment, such as a hood or a glove box.
  - 5.1.3. Assembly and disassembly of glassware creates a risk of breakage and cuts. All staff members shall wear Kevlar or MAPA blue latex cut-resistant gloves over chemically resistant gloves when assembling and disassembling glassware.
  - 5.1.4. Eye protection that satisfies ANSI Z87.1, laboratory coat, and chemically resistant gloves must be worn while samples, standards, solvents, and reagents are being handled. Latex and vinyl gloves provide no protection against most of the organic solvents used in this method. Nitrile or similar gloves must be used. Latex gloves may be used for methanol.
  - 5.1.5. Exposure to chemicals must be maintained as low as reasonably achievable, therefore all samples must be opened, transferred and prepared in a fume hood. Solvent and waste containers will be kept closed unless transfers are being made.
  - 5.1.6. Laboratory procedures such as repetitive use of pipets, repetitive transferring of extracts, and manipulation of filled separatory funnels and other glassware represent a significant potential for repetitive motion or other ergonomic injuries. Laboratory associates performing these procedures are in the best position to realize when they are at risk for these types of injuries. Whenever a situation is found in which an employee is performing the same repetitive motion, the employee shall immediately bring this to the attention of their supervisor, manager, or the EH&S staff. The task will be analyzed to determine a better means of accomplishing it.

5.1.7. The use of separatory funnels to extract aqueous samples with methylene chloride creates excessive pressure very rapidly. The use of separatory funnels during the partition and back extraction of sample extracts can also create excessive pressure. Initial venting should be done immediately after the sample container has been sealed and inverted. Vent the funnel into the hood away from people and other samples. This is considered a high-risk activity, and a face shield must be worn over safety glasses or goggles when it is performed. Alternately, the extraction can be performed behind a closed fume hood sash on a mechanical shaker.

5.1.8. When Dean-Stark/Soxhlet clean-ups or extractions are performed overnight or unattended, special precautions must be taken. Open the chiller valves to the system about 15 minutes before the heating elements are turned on, and check every condenser to ensure that it is cold and functioning properly before turning the heating elements on. Check every condenser again about 15 minutes after turning the heating elements on to ensure that they are still cold and functioning properly. If the system is left operating overnight or unattended for an extended period, the first chemist to come back into the lab must again check every condenser to ensure that it is still cold and functioning properly.

# 5.2. Primary Materials Used

The following is a list of the materials used in this method, which have a serious or significant hazard rating. **NOTE:** This list does not include all materials used in the method. The table contains a summary of the primary hazards listed in the MSDS for each of the materials listed in the table. A complete list of materials used in the method can be found in the reagents and materials section. Employees must review the information in the MSDS for each material before using it for the first time or when there are major changes to the MSDS.

Material (1)	Hazards	Exposure Limit (2)	Signs and symptoms of exposure
Acetone	Flammable	1000 ppm- TWA	Inhalation of vapors irritates the respiratory tract. May cause coughing, dizziness, dullness, and headache.
Benzene	Flammable Toxic Carcinogen	PEL: 1 ppm TWA; 5 ppm 15 MIN. STEL	Causes skin irritation. Toxic if absorbed through skin. Causes severe eye irritation. Toxic if inhaled. Vapor or mist causes irritation to mucous membranes and upper respiratory tract. Exposure can cause narcotic effect. Inhalation at high concentrations may have an initial stimulatory effect on the central nervous system characterized by exhilaration, nervous excitation and/or giddiness, depression, drowsiness or fatigue. Victim may experience tightness in the chest, breathlessness, and loss of consciousness.
Cyclohexane	Flammable Irritant	300 ppm TWA	Inhalation of vapors causes irritation to the respiratory tract. Symptoms may include coughing, shortness of breath. High concentrations have a narcotic effect.

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Material (1)	Hazards	Exposure Limit (2)	Signs and symptoms of exposure		
Hexane	Flammable Irritant	500 ppm-TWA	Inhalation of vapors irritates the respiratory tract. Overexposure may cause lightheadedness, nausea, headache, and blurred vision. Vapors may cause irritation to the skin and eyes.		
Isooctane	Flammable Irritant	None established	Inhalation of vapors may cause nausea, headache, dizziness, loss of consciousness, irritation to upper respiratory tract, pain in throat and nose, coughing, wheezing, shortness of breath.		
Methanol	Flammable Poison Irritant	200 ppm-TWA	A slight irritant to the mucous membranes. Toxic effects exerted upon nervous system, particularly the optic nerve. Symptoms of overexposure may include headache, drowsiness and dizziness. Methyl alcohol is a defatting agent and may cause skin to become dry and cracked. Skin absorption can occur; symptoms may parallel inhalation exposure. Irritant to the eyes.		
Methylene Chloride	Carcinogen Irritant	25 ppm-TWA 125 ppm-STEL	Causes irritation to respiratory tract. Has a strong narcotic effect with symptoms of mental confusion, light-headedness, fatigue, nausea, vomiting and headache. Causes irritation, redness and pain to the skin and eyes. Prolonged contact can cause burns. Liquid degreases the skin. May be absorbed through skin.		
Sodium Hydroxide	Corrosive Poison	2 ppm, 5 mg/m <sup>3</sup>	This material will cause burns if comes into contact with the skin or eyes. Inhalation of Sodium Hydroxide dust will cause irritation of the nasal and respiratory system.		
Sulfuric Acid (1)	Corrosive Oxidizer Dehydra-dator	1 mg/m <sup>3</sup>	This material will cause burns if comes into contact with the skin or eyes. Inhalation of vapors will cause irritation of the nasal and respiratory system.		
Tetradecane	Irritant	None established	Inhalation of vapors may cause difficulty breathing, headache, intoxication and central nervous system damage.		
Toluene	Flammable Poison Irritant	200 ppm-TWA 300 ppm- Ceiling	Inhalation may cause irritation of the upper respiratory tract. Symptoms of overexposure may include fatigue, confusion, headache, dizziness and drowsiness. Peculiar skin sensations (e. g. pins and needles) or numbness may be produced. Causes severe eye and skin irritation with redness and pain. May be absorbed through the skin.		
	1 – Always add acid to water to prevent violent reactions.				
2 – Exposure	e limit refers to th	ne OSHA regulat	ory exposure limit.		

# **6. EQUIPMENT AND SUPPLIES**

The following list of items does not necessarily constitute an exhaustive compendium of the equipment needed for this analytical method.

- 6.1. Nitrogen evaporation apparatus with variable flow rate.
- 6.2. Balances capable of accurately weighing to 0.01 g and 0.0001 g.
- 6.3. Centrifuge.
- 6.4. Water bath, equipped with concentric ring covers and capable of maintaining temperature control within  $\pm 2^{\circ}$ C.

- 6.5. Stainless steel or glass containers large enough to hold contents of one-pint sample containers.
- 6.6. Drying oven.
- 6.7. Stainless steel spoons and spatulas.
- 6.8. Pipettes, disposable, Pasteur, 150 mm long x 5 mm ID.
- 6.9. Pipettes, disposable, serological, 10 mL, for the preparation of the carbon column specified in Section 7.1.
- 6.10. Reacti-vial, 2 mL, silanized clear glass.
- 6.11. Stainless steel meat grinder with a 3- to 5-mm hole size inner plate.
- 6.12. Separatory funnels, 250 mL.
- 6.13. Separatory funnels, 1000 mL.
- 6.14. Teflon® boiling chips (or equivalent) washed with methylene chloride before use.
- 6.15. Chromatographic column, glass, 300 mm x 10.5 mm, fitted with Teflon® stopcock.
- 6.16. Adapters for concentrator tubes.
- 6.17. Glass fiber filters, Whatman GF-D, GF-F, GMF150, or equivalent.
- 6.18. Solid phase extraction discs, 3M 90mm C18, or equivalent.
- 6.19. Dean-Stark trap, 5 or 10 mL, with T-joints, condenser and 125 mL flask.
- 6.20. Continuous liquid-liquid extractor.
- 6.21. All-glass Soxhlet apparatus, 500 mL flask.
- 6.22. Soxtherm extraction apparatus (or equivalent), including glass thimble holders, glass beakers, and gaskets.
- 6.23. Glass funnels, sized to hold 170 mL of liquid.
- 6.24. Desiccator.
- 6.25. Turbo evaporator
- 6.26. Rotary evaporator with a temperature controlled water bath.

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- 6.27. High speed tissue homogenizer, equipped with an EN-8 probe or equivalent.
- 6.28. Glass wool, extracted with methylene chloride, dried and stored in a clean glass jar.
- 6.29. Vacuum extraction device for solid phase extraction, 1 Liter glass funnel with 90mm filter disc holder with a vacuum source, Kontes or equivalent.

## 7. REAGENTS AND STANDARDS

- 7.1. Column Chromatography Reagents
  - 7.1.1. Silica Gel Kieselgel 60 or equivalent, activate for 1 hour at 184°C before use. Store at 130°C in covered flask.
  - 7.1.2. Acid Alumina ICN or equivalent, activated as necessary.
  - 7.1.3. Basic Alumina ICN or equivalent. No activation required.
  - 7.1.4. Granular carbon/silica gel Mix 3.6 g granular carbon and 16.4 g activated silica gel; (alternatively, prepare carbon/silica gel (5%/95%); i.e., combine 5 g precleaned carbon with 95 g silica gel). Store at room temperature in a Teflon ® lined covered jar. The first LCS prepared with a new batch of column packing material is the quality control check of the packing materials. Refer to historical control limits before accepting the new batch of material.
  - 7.1.5. 44% H<sub>2</sub>SO<sub>4</sub> /silica gel Mix 24 mL conc. H<sub>2</sub>SO<sub>4</sub> and 56 g activated silica gel. Stir and shake until free flowing. Store at room temperature.
  - 7.1.6. 33% NaOH/silica gel Mix 34 mL 1N NaOH and 67 g activated silica gel. Stir and shake until free flowing. Store at room temperature.

## 7.2. Acid Alumina Activity Assessment

Alumina activity may vary with the matrix or environmental conditions. Monitor internal standard and cleanup recovery standard recoveries in extract analysis. Low recoveries of cleanup recovery standard (CRS) may indicate loss of alumina activity. Assess stability of alumina activity and apply corrective action as appropriate (reactivate and reprofile).

Note: a column profile should be done to show elution of all 2,3,7,8 substituted analogs so problems can be readily identified.

7.2.1. Profile each vendor lot of activated alumina as corrective action for low internal standard and CRS recoveries dictate. If necessary, proceed as follows:

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- 7.2.1.1. Set up and label 3 acid alumina columns.
- 7.2.1.2. Pre-rinse with 20 mL hexane.
- 7.2.1.3. Add 2 mL hexane spiked with internal standards and natives (spike amounts equivalent to those for LCS) with 2X2 mL hexane rinse of fractions.
- 7.2.1.4. Elute each column with 20 mL hexane. Collect and label these fractions
- 7.2.1.5. Elute each column with 5 x10 mL methylene chloride/hexane at the appropriate v/v percent. Collect and label these fractions separately.
- 7.2.1.6. Elute each column with 10 mL of 100% methylene chloride. Collect and label these fractions. Reduce all fractions to final volume and add recovery standard.
- 7.2.2. Review data and select an elution scheme. Group the fraction from each solvent system as follows:
  - 7.2.2.1. Pre-analyte fraction consists of all eluent prior to elution of first target analytes.
  - 7.2.2.2. Analyte fraction consists of all that contain detectable levels of target analytes.
  - 7.2.2.3. Post-analyte fraction consists of all eluents after elution of the last target analyte.
- 7.2.3. Select the solvent system which best meets the following two conditions:
  - 7.2.3.1. Pre-analyte fraction consists of 20mL hexane and no more than 20 mL mixed solvent.
  - 7.2.3.2. Analyte fraction consists of no more than 20mL of mixed solvent and contains greater than 90% of all target analytes and greater than 80% of all internal standards.
- 7.2.4. After selection of the appropriate solvent system and fractionation pattern, perform triplicate acid alumina cleanups on spiked hexane to ensure reproducibility of the fractionation pattern. Document each elution scheme.
- 7.2.5. Each subsequent batch of acid alumina used in the lab (from the same vendor lot) must be checked for stable activity.

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# 7.3. Reagents

- 7.3.1. Sulfuric acid, concentrated, ACS grade, specific gravity 1.84.
- 7.3.2. Distilled water demonstrated to be free of interferents
- 7.3.3. 1 N HCl.
- 7.3.4. Silica gel.
- 7.3.5. Solution for breaking emulsions: Slowly add 1.0L of reagent grade NaOH solution to a 2.0L NaOH container, containing 1.0L of DI H2O, and leave the container in secondary containment with the lid off.

# Warning: The solution will begin to heat so let the solution stand until equilibrium is met and the solution is at room temperature.

When this process is complete, the solution will then be ready for use in the samples.

- 7.3.6. Precleaned Sodium Sulfate.
- 7.3.7. Canola Oil (for tissue extraction only), or other suitable oil.
- 7.4. Desiccating Agent
  - 7.4.1. Sodium sulfate, granular, anhydrous.
- 7.5. Solvents
  - 7.5.1. High-purity, distilled-in-glass or highest available purity: Methylene chloride, hexane, methanol, tetradecane, isooctane, toluene, cyclohexane, and acetone.
- 7.6. All daily internal standard, daily clean up recovery standards, and daily spiking solutions are stable for one year from preparation. After 1 year, solutions may be reverified. The re-verified solution may be used for an additional year, or until there is evidence of compound degradation or concentration. The re-verification must be performed using an unexpired, not previously re-verified solution from a second lot or second yendor.
  - 7.6.1. Sealed ampules may be used until the manufacturer's expiration date is exceeded. If no expiration date is provided, then the expiration date will be 10 years from the date the ampule is opened. The solvent level should be monitored prior to each use to assure there has been no concentration of the standard over time.

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7.6.2. Standards for method 8290A require storage at  $\leq 6^{\circ}$ C.

# 7.7. Field Surrogate Solution (air matrices)

This solution contains one <sup>37</sup>Cl labeled analog (for Method TO-9/TO-9A) or one <sup>37</sup>Cl and four <sup>13</sup>C labeled analogs (for Method 0023) at the nominal concentration indicated in Table 2. It is used to assess sample collection and recovery procedures.

## 7.8. Internal Standard

This isooctane solution contains the nine internal standards at the nominal concentrations that are listed in Table 2. The solution contains at least one carbon-labeled standard for each homologous series, and it is used to measure the concentrations of the native substances. (Note that  $^{13}C_{12}$  -OCDF is not present in the solution.)

# 7.9. Native Spike Standard

Also known as the Matrix Spike or Native Spike solution. Contains all the 2,3,7,8-substituted unlabeled analytes listed in Table 2. Prepare using the appropriate standards to yield a spiking solution with a concentration of 4.0 ng/ml for the tetra-CDDs/CDFs, 20 ng/ml for the penta-, hexa-, and hepta- CDDs/CDFs, and 40 ng/ml for the octa- CDD/CDF.

# 7.10. Recovery Standard Solution

This tetradecane solution contains two recovery standards ( ${}^{13}C_{12}$ -1,2,3,4-TCDD and  ${}^{13}C_{12}$ -1,2,3,7,8,HxCDD). An appropriate volume of this solution is spiked into each sample extract before the final concentration step.

# 7.11. Cleanup Recovery Standard Solution (CRS)

Prepare <sup>37</sup>Cl<sub>4</sub>-2,3,7,8-TCDD at the concentration shown in Table 2, in isooctane (or toluene).

# 7.12. Preparation and QC of PUF material

- 7.12.1. The PUF material is purchased pre-cut.
- 7.12.2. The PUFs are rinsed by Soxhlet with acetone (or other appropriate solvent) for a minimum of 16 hours and air dried for a minimum of 2 hours in a contaminant-free area.
- 7.12.3. One PUF from the rinsed batch is randomly selected to be the QC sample for the batch.
- 7.12.4. The PUF is loaded into a pre-cleaned Soxhlet extractor charged with toluene.
- 7.12.5. The 1613/8290 daily internal standard solution is spiked into the PUF and it

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is extracted for a minimum of 16 hours.

- 7.12.6. The Soxhlet extract is recovered and processed according to Section 11.4.
- 7.12.7. The batch of PUF is considered acceptable if no target analytes are detected at or above the laboratory or project specific reporting limit.

# 8. SAMPLE COLLECTION, PRESERVATION AND STORAGE

- 8.1. The sample collection, shipping, handling, and chain-of-custody procedures are not described in this document. Sample collection personnel will, to the extent possible, homogenize samples in the field before filling the sample containers. This should minimize or eliminate the necessity for sample homogenization in the laboratory. The analyst should make a judgment, based on the appearance of the sample, regarding the necessity for additional mixing. If the sample is clearly non-homogeneous, the entire contents should be transferred to a glass or stainless steel pan for mixing with a stainless steel spoon or spatula before removal of a sample portion for analysis.
- 8.2. Grab and composite samples must be collected in glass containers.
- 8.3. Ambient air samples are collected on a Quartz Fiber Filter followed by a glass sleeve containing a polyurethane foam plug.
- 8.4. Samples from stationary sources are collected on glass or quartz fiber filters and XAD-2 Resin. (See WS-ID-0009 for sample preparation procedures).
- 8.5. Conventional sampling practices must be followed. Do not rinse the bottle with sample before collection. Sampling equipment must be free of potential sources of contamination.
- 8.6. Grinding or blending of fish samples.

If not otherwise specified by the client, the whole fish (frozen) should be blended or ground to provide a homogeneous sample. The use of a stainless steel meat grinder with a 3 to 5 mm hole size inner plate is recommended. In some circumstances, analysis of fillet or specific organs of fish may be requested by the client. If so requested by the client, the above whole fish requirement is superseded. More detail can be found in "Tissue Sampling and Handling for a variety of Methods" (WS-WI-0018).

# Warning: Hearing protection must be worn when grinding samples.

8.7. With the exception of the fish tissues, which must be stored at -  $20^{\circ}$ C, all samples should be stored at  $4^{\circ}$ C  $\pm$  2, extracted within 30 days and completely analyzed within 45 days of collection. The 30 day hold time is recommended. PCDDs and PCDFs have demonstrated stability for greater than one year.

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8.8. All extracts must be stored capped, in the dark, at room temperature (approximately 21°C to 28°C). All extracts for method 8290A must be stored capped at  $\leq$  6°C.

8.9. For moisture determinations refer to SOP WS-OP-0013.

## 9. **QUALITY CONTROL**

9.1. One method blank (MB) must be extracted with every process batch of similar matrix, not to exceed twenty (20) samples. The method blank is an aliquot of laboratory matrix (reagent water, sodium sulfate, PUF, XAD, filter, etc.) processed in the same manner and at the same time as the associated samples. Corrective actions must be documented on a Non-Conformance memo, then implemented when target analytes are detected in the method blank above the reporting limit or when surrogate recoveries are outside control limits. Re-extraction of the blank, other batch QC, and the affected samples are required when the method blank is deemed unacceptable. The method blank contains a PUF plug, XAD, or filter prepared from the same batch as the field samples whenever possible for air samples.

Certain programs, such as DOD, may require a more stringent evaluation of the method blank, for instance, that the blank not contain any analytes of interest at a concentration greater than ½ the lower calibration limit.

Note: Re-extraction of the blank, QC and affected samples for the air matrices (PUF, XAD, and filter) is not generally possible because the entire sample is consumed in the initial extraction.

- 9.1.1. If the accompanying samples are aqueous, use distilled water as a matrix. Take the method blank through all steps detailed in the analytical procedure.
- 9.1.2. Use sodium sulfate as the method laboratory matrix when solids are extracted. Use a mixture of sodium sulfate and canola oil as the matrix when tissues are extracted. Take the method blank through all steps detailed in the analytical procedure.
- 9.1.3. The method blank must be spiked prior to extraction with the same amount of <sup>13</sup>C -labeled internal standards as added to samples.
- 9.1.4. If method blank contamination is present, check solvents, reagents, fortification solutions, apparatus and glassware to locate and eliminate the source of contamination before any further samples are extracted and analyzed. The presence of any analyte in the method blank ate concentrations greater than the reporting limit (RL) is cause for corrective action.
  - 9.1.4.1. OCDD is a ubiquitous laboratory contaminant. A method blank and the associated samples are deemed acceptable if the OCDD

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- concentration is <5x the specified reporting limit. Flag data appropriately. The analyst is expected to investigate and eliminate potential sources of systematic contamination.
- 9.1.4.2. If a target analyte is detected in the blank but the associated samples are ND (not detected), then the data may be reported, unless otherwise directed by the client. Note the action in the narrative
- 9.1.4.3. If a target analyte is detected in the blank, but the concentration of the contaminant in the samples >10x the blank concentration, then the data may be reported, unless otherwise directed by the client. Note the action in the narrative.
- 9.1.4.4. If one of the conditions above is not met then the sample associated with a contaminated method blank must be reextracted
- 9.1.5. If new batches of reagents or solvents contain interfering contaminants, purify or discard them.
- 9.2. A Laboratory Control Sample (LCS) must be extracted with every process batch of similar matrix, not to exceed twenty (20) samples. The LCS is an aliquot of laboratory matrix (e.g. water, sodium sulfate, PUF, XAD, etc.) spiked with analytes of known identity and concentration. The LCS must be processed in the same manner and at the same time as the associated samples. Corrective actions must be documented on a Non-Conformance memo, then implemented when recoveries of any spiked analyte is outside control limits provided on the LIMS or by the client. Re-extraction of the blank, other batch QC and all associated samples are required if the LCS is deemed unacceptable. See policy WS-PQA-003 for specific acceptance criteria. When associated with PUF samples, the LCS should contain a PUF plug prepared from the same batch as the field samples whenever possible.

Note: Re-extraction of the blank, QC and affected samples for the air matrices (PUF, XAD, and filter) is not generally possible because the entire sample is consumed in the initial extraction

- 9.2.1. A LCS is deemed acceptable if control analytes are above control limits and the associated samples are ND, unless otherwise specified by the client. Note any actions in the narrative.
- 9.3. The assessment of matrix effects on method performance, as required by NELAP, is met in Method 8290 and 8290A, as in all isotope dilution techniques, with the use of isotopically labeled compounds. These isotopically labeled compounds are analogs of target analytes and are spiked into each sample. Therefore, matrix effects on method performance may be judged by the recovery of these analogs. Sample analysis

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acceptance is controlled by the performance of these analogs in each sample. A Matrix Spike/Matrix Spike Duplicate (MS/MSD or MS/SD) pair are extracted at the client's request only. Method 8290A does not address analysis of MS/MSD. An exception to this rule is a batch containing South Carolina samples for Method 8290. These batches must have an MS/MSD prepared. However, South Carolina requires Method 8290A after December 31, 2008. An MS/MSD pair are aliquots of a selected field sample spiked with analytes of known identity and concentration. When requested by the client, the MS/MSD pair shall be processed in the same manner and at the same time as the associated samples. Corrective actions must be documented on a Non-Conformance memo, then implemented when recoveries of any spike analyte is outside control limits provided on the LIMS or by the client. Re-extraction of the blank, the LCS, the selected field sample, and the MS/MSD may be required after evaluation and review. Matrix Spike/ Matrix Spike Duplicates are not generally applicable for air samples due to the difficulty in collecting identical or representative samples. An LCS/LCSD may be extracted to show precision of the extraction and analysis process.

- 9.3.1. Matrix Spike (MS): A sample, which is spiked with a known amount of the matrix spike fortification solution prior to the extraction step. The recoveries of the matrix spike compounds are determined; they are used to estimate the effect of the sample matrix upon the analytical methodology.
- 9.3.2. Matrix Spike Duplicate (MSD): A second portion of the same sample as used in the matrix spike analysis and which is treated like the matrix spike sample.
- 9.3.3. Locate the sample for the MS and MSD analyses (the sample may be labeled "double volume").
- 9.3.4. Add an appropriate volume of the matrix spike fortification solution, adjusting the fortification level as specified in Table 1, under IS Spiking Levels.
- 9.3.5. The results obtained from the MS and MSD samples (percent recovery and concentrations of 2,3,7,8-substituted PCDDs/PCDFs) should agree within 20 percent relative difference. Report all results and flag outliers.
- 9.3.6. Internal standard recoveries are flagged if they are outside the recovery goals. Re-extraction of affected samples should be performed if signal-to-noise for any internal standard is less than 10:1.

# 9.4. Duplicates

9.4.1. Upon client request, duplicates may be processed. Locate the sample specified for duplicate analysis, and prepare and analyze a second 10-g soil or sediment sample portion or 1 L water sample, or an appropriate amount of

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the type of matrix under consideration. Duplicate samples are not generally applicable for air samples due to the difficulty in collecting identical or representative samples. A duplicate injection of a sample extract may be performed to display instrument precision.

- 9.4.1.1. The results of the laboratory duplicates (percent recovery and concentrations of 2,3,7,8-substituted PCDD/PCDF compounds) should agree within 25 percent relative difference. Report all results and flag outliers.
- 9.4.2. Internal standard recoveries are flagged if they are outside the recovery goals. Re-extraction of affected samples should be performed if signal-to-noise for any internal standard is less than 10:1.

#### 9.5 Field Blanks

- 9.5.1. Each batch of samples may contain a field blank sample of nominally uncontaminated soil, sediment or water that is to be processed for analysis.
  - 9.5.1.1. Weigh a 10-g portion or use 1 L (for aqueous samples) of the specified field blank sample and add the appropriate amount of internal standard to yield 100 pg/ $\mu$ L in the final extract.
  - 9.5.1.2. Extract by using the procedures described in Section 11. As applicable, add the appropriate amount of recovery standard to yield 100 pg/ $\mu$ L in the final extract. Analyze a 1-2  $\mu$ L aliquot of the concentrated extract using SOP WS-ID-0005.

# 9.6. Rinsate Samples

- 9.6.1. In addition to the field blank, a batch of samples may include a rinsate, which is a portion of the solvent (usually trichloroethylene) that was used to rinse sampling equipment. The rinsate is analyzed to assure that the samples were not contaminated by the sampling equipment.
- 9.6.2. The rinsate sample must be processed like a regular sample.

  Take a 100-mL (± 0.5 mL) portion of the sampling equipment rinse solvent (rinsate sample), filter, if necessary, and add the appropriate amount of internal standard to yield 100 pg/μL in the final extract.
- 9.6.3. Using appropriate methods, concentrate to approximately 10 mL.
- 9.6.4. Just before analysis, add the appropriate amount of recovery standard to yield 100 pg/ $\mu$ L in the final extract. Reduce the volume to a final volume of 20  $\mu$ L, as necessary. No column chromatography is required.

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9.6.5. Analyze an aliquot following the same procedures used to analyze samples.

# 9.7. Surrogate/Clean Up Recovery Standard

A surrogate compound may be spiked into all air media samples prior to collection. For all other matrices, a clean up recovery standard is spiked following extraction and just prior to cleanup, in order to monitor relative loss of internal standard during both extraction and cleanup.

#### 9.8. Internal Standards

An internal standard is a <sup>13</sup>C -labeled analog of a PCDD/PCDF congener. Internal standards are added to all samples including method blanks and quality control samples before extraction, and they are used to quantitate the concentration of the analytes. Nine internal standards are used in this method. There is one for each of the dioxin and furan homologs (except for OCDF) with the degree of chlorination ranging from four to eight. Additional internal standards may be added to act as retention time references, but they are not used for quantitation.

- 9.8.1. A 2000 pg aliquot of the internal standard mixture is added to all samples, regardless of sample size. As an example, for <sup>13</sup>C<sub>12</sub> -2,3,7,8-TCDD, a 10-g soil sample requires the addition of 2000 pg of <sup>13</sup>C<sub>12</sub> -2,3,7,8-TCDD to give the requisite fortification level.
- 9.8.2. Internal standards must be spiked into all samples, QC samples, and included in all calibrations.
- 9.8.3. For each sample and QC aliquot, calculate the percent recovery. The percent recovery should be between 40 percent and 135 percent for all nine internal standards.
- 9.8.4. A low or high percent recovery for a blank does not require discarding the analytical data but it may indicate a potential problem with future analytical data. Internal standard recoveries are flagged if they are outside the recovery goals. Re-extraction of affected samples should be performed if signal-to-noise for any internal standard is less than 10:1.
- 9.9. Recovery Standard: Two recovery standards are used to determine the percent recoveries for the internal standards. The <sup>13</sup>C<sub>12</sub> -1,2,3,4-TCDD is used to measure the percent recoveries of the tetra- and pentachlorinated internal standards while <sup>13</sup>C<sub>12</sub> -1,2,3,7,8,9-HxCDD is used to determine the recovery of the hexa-hepta- and octachlorinated internal standards. <sup>13</sup>C<sub>12</sub> -1,2,3,7,8,9-HxCDD also acts as a retention time reference for the unlabeled analog present in sample extracts. They are added to the final sample extract before HRGC/HRMS instrument analysis.
- 9.10. Recommended Corrective Actions and Troubleshooting Steps

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- Verify satisfactory instrument performance.
- If possible, verify that no error was made while weighing the sample aliquots.
- Review the analytical procedures with the performing laboratory personnel.

#### 10. CALIBRATION

- 10.1. On a daily basis, calibrate any balance to be used in accordance with SOP WS-QA-0041.
- 10.2. On a monthly basis, calibrate any autopipettor to be used in accordance with SOP WS-QA-0004.

#### 11. PROCEDURE

- 11.1. One time procedural variations are allowed only if deemed necessary in the professional judgment of the supervisor to accommodate variation in sample matrix, radioactivity, chemistry, sample size, or other parameters. Any variation in procedure shall be completely documented using a Nonconformance Memo and is approved by a Technical Specialist and QA Manager. If contractually required, the client shall be notified. The Nonconformance Memo shall be filed in the project file.

  Any unauthorized deviations from this procedure must also be documented as a nonconformance, with a cause and corrective action described.
- 11.2. Refer to SOP WS-ID-0009 for the preparation of stationary source samples.
- 11.3. Sample Pre-Treatment
  - 11.3.1. Paper Pulp Sludges are generally air-dried and ground prior to extraction following Section 11.5. Because of the drying procedure, a Dean-Stark water separator is optional for extraction.
  - 11.3.2. Fly Ash Fly ash samples are pretreated with HCl prior to extraction by both soxhlet and separatory funnel techniques.
    - 11.3.2.1. Weigh 2-10g of sample aliquot into a clean glass jar.
    - 11.3.2.2. Add 1.0mL of the internal standard mixture with 2 mL of acetone.
    - 11.3.2.3. Add 150 mL of 1N hydrochloric acid and shake for 4 hours.
    - 11.3.2.4. If the sample reacts violently with acid, then allow the sample to equilibrate for 4 hours with no shaking.
    - 11.3.2.5. Filter the contents of the jar through a glass fiber filter.

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- 11.3.2.6. Extract the solids as per Section 11.5, omitting the daily internal standard spike for the samples.
- 11.3.2.7. Extract the aqueous filtrate as per Section 11.8, using 100 mL of toluene for the first shake, and 100 mL of hexane for subsequent shakes.
- 11.3.2.8. Concentrate the combined toluene solutions to near dryness on a rotary evaporator at 50°C. Proceed with Section 11.12 as necessary.

Note: As an option, a Soxhlet/Dean Stark extractor system may be used, with toluene as the solvent. No sodium sulfate is added when using this option.

- 11.4. Waste Dilution (Still-Bottom/Fuel Oil, and other solvent-miscible materials).
  - 11.4.1. Weigh 1 g of the waste (organic liquids, fuel oils, and solids that will dissolve in a solvent) into a vial.
  - 11.4.2. Add 40 mL of toluene (or other solvent if the material is not miscible/soluble in toluene). Shake gently to dissolve.
  - 11.4.3. Remove a 4.0 mL aliquot (0.1g sample equivalent) and place in a culture tube. Add 1.0 mL of daily internal standard and 1.0 mL of cleanup recovery standard, and proceed to Section 11.12.
- 11.5. Soxhlet Extraction (Solids, Tissues, Sludges, Wipes)
  - 11.5.1. Pre-extract the glassware by heating the flask until the toluene is boiling. When properly adjusted, 1-2 drops of toluene per second will fall from the condenser tip into the receiver. Extract the apparatus for a minimum of four hours.

WARNING: Open the chiller supply valves about 15 minutes before turning on the heating element and ensure that all of the condensors are cold before you turn the heating element on. Check all of the condensors about 15 minutes after starting the heating process to ensure that they are still cold and functioning properly. If this cleaning cycle is to be left unattended (e.g., overnight) the first chemist to arrive the next morning is to check all condensers to ensure that they are still cold and functioning properly.

- 11.5.2. After pre-extraction, cool and disassemble the apparatus.
- 11.5.3. If tissues requiring % Lipids are to be extracted, for each sample weigh the concentration vessel with label and boiling chips. Record the mass on the benchsheet. Refer to SOP WS-QA-0018 "Subsampling", for instructions on how to homogenize and subsample the container of sample.

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- 11.5.4. Weigh a well-mixed aliquot of each sample (10 g, unless otherwise specified) into a clean Soxhlet thimble. Record the mass to the nearest 0.01g. Use sodium sulfate for the batch QC (MB, LCS) for solids, and a mixture of 9 g sodium sulfate and 1 g canola oil for the batch QC for tissue matrices.
  - 11.5.4.1. In the case of wipes, place the entire wipe sample into the Soxhlet apparatus (no thimble needed), including any liquid present with the sample. Use pre-cleaned wipes for the batch QC samples.
- 11.5.5. Place the thimble into a Soxhlet apparatus equipped with a Dean-Stark water separator.
- 11.5.6. Spike all samples with 1.0 mL of internal standard solution (2 pg/ $\mu$ L), for a final concentration of 200 pg/g (based on a 10 g sample).
- 11.5.7. Spike the LCS (and MS/MSD, if present) with 50 uL of native spike.
- 11.5.8. Reassemble the pre-extracted apparatus and add a fresh charge (250-300 mL) of toluene to the receiver and reflux flask.
- 11.5.9. Reflux 16 hours, with the solvent cycling at least 5 times per hour.

WARNING: Open the chiller supply valves about 15 minutes before turning on the heating element and ensure that all of the condensors are cold before you turn the heating element on. Check all of the condensors about 15 minutes after starting the heating process to ensure that they are still cold and functioning properly. If this cleaning cycle is to be left unattended (e.g., overnight) the first chemist to arrive the next morning is to check all condensers to ensure that they are still cold and functioning properly.

11.5.10. Drain the water from the receiver if the receiver fills with water. Check and drain when necessary.

Note: If the receiver holds 10 mL of liquid, and 20 g of an approximately 10% solid sample is being extracted, then approximately 9 mL of water will end up in the receiver. In this case, the receiver will not need to be emptied (insufficient liquid to overflow), but it should be checked. If the sample amount is 50, and the percent solids is still 10%, then 45 mL of water will end up in the receiver. In this case, frequent checking is required, and the receiver will need to be emptied at least 5 times.

- 11.5.11. After refluxing, allow the apparatus to cool.
- 11.5.12. If samples DO NOT require % lipids add 100 μL of tetradecane as a keeper to the round bottom flask.

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- 11.5.13. Proceed to Section 11.17.
- 11.6. SoxTherm Extraction (Solids, Tissues, Sludges, Wipes)
  - 11.6.1. Prior to loading samples, run the system through 2 cleaning cycles (approximately 1 hour each).
  - 11.6.2. After pre-extraction, cool and disassemble the apparatus.
  - 11.6.3. Weigh a well-mixed aliquot of each sample (10 g, unless otherwise specified) into a clean Soxhlet thimble. Record the mass to the nearest 0.01g. Use sodium sulfate for the batch QC (MB, LCS) for solids, and a mixture of 9 g sodium sulfate and 1 g canola oil for the batch QC for tissue matrices.
    - 11.6.3.1. In the case of wipes, place the entire wipe sample into the Soxhlet apparatus (no thimble needed), including any liquid present with the sample. Use pre-cleaned wipes for the batch QC samples.
  - 11.6.4. Place the thimble into the Soxtherm apparatus.
  - 11.6.5. Spike all samples with 1.0 mL of internal standard solution (2 pg/μL), for a final concentration of 200 pg/g (based on a 10 g sample).
  - 11.6.6. Spike the LCS (and MS/MSD, if present) with 50 uL of native spike.
  - 11.6.7. Reassemble the pre-extracted apparatus and add a fresh charge (150 mL) of toluene to the apparatus.
  - 11.6.8. Program the system to boil for 1 hour, and reduce the toluene volume by 70-90 mL (volume < volume of the thimble).
  - 11.6.9. Continue the extraction for one hour fifteen minutes, reducing the toluene volume by another 15 mL.
  - 11.6.10. After refluxing, allow the apparatus to cool.
  - 11.6.11. Pour the samples into round bottom flasks, and if samples DO NOT require % lipids add 100  $\mu$ L of tetradecane as a keeper to the round bottom flask.
  - 11.6.12. Proceed to Section 11.17.
- 11.7. Extract Splitting (Wipes)

Wipe extracts prepared using either Soxhlet or shaking techniques are split prior to further workup, to permit an archive aliquot, or analysis by an additional method.

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Once the extract has been concentrated using the rotovap or Turbovap, proceed as follows:

- 11.7.1. Add approximately 1 mL of hexane or toluene to rinse the sides of the round bottom flask. Using a pipette, withdraw the sample from the round bottom flask and transfer the liquid into a test-tube. Use additional amounts of solvents to rinse the flask. Transfer all the liquid into the test-tube. Ensure that all traces of sample in the round bottom flask have been thoroughly rinsed from all surfaces. Bring the sample volume to 8.0 mL or 10.0 mL (or appropriate volume) with the addition of rinse solvent.
- 11.7.2. Upon completion of the rinsing, cap the test tube and shake vigorously. Take ½ of each sample (or an appropriate amount as instructed by the client, program manager or department manager) and transfer to a culture tube. Archive the remaining sample for future use.
  - 11.7.2.1. If only one analysis is required, then ½ of the sample is archived and the other half is analyzed.
  - 11.7.2.2. If "N" analyses are required, then the extract is divided into "N+1" equal portions, so that one portion is archived, and a portion is used for each test.
- 11.8. Aqueous Samples (liquid/liquid extraction).
  - 11.8.1. When setting up the glassware for a batch, for each sample label one separatory funnel and one 500 mL round-bottom flask with the sample ID.
  - 11.8.2. Weigh the sample in the bottle on the top loading balance to the nearest centigram (0.01g), and record the mass.
  - 11.8.3. For each sample, add 1 mL of daily internal standard solution into 2 mL of acetone. Add this solution to the sample in the separatory funnel. Each aliquot of spike mixture is added similarly.
  - 11.8.4. Dissolve 50µL of the target analyte into acetone and add this mixture into the LCS container.
  - 11.8.5. Pour the entire sample (approximately 1L) into a 2L separatory funnel that is labeled with the sample ID.
  - 11.8.6. Add 100 mL methylene chloride to the sample bottle, seal, and shake for 30 seconds to rinse the inner surface. Transfer the solvent to the separatory funnel.
  - 11.8.7. Create a blank and LCS by adding 1 L of laboratory reagent water to 2

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- additional separatory funnels. Add 100 mL methylene choride to each funnel.
- 11.8.8. To the LCS, add 50 μL of the precision and recovery standard dissolved into 2 mL of acetone.
- 11.8.9. Extract the samples by shaking each funnel for two minutes with periodic venting.

Warning: Separatory funnel extraction with methylene chloride is a high-risk activity. Pressure may build rapidly in the funnel. It should be vented after several seconds of shaking, and often enough to prevent build-up of pressure. Chemist performing separatory funnel extraction must wear a face shield over their safety glasses/goggles. Alternatively, the extraction can be performed behind a closed fume hood sash.

- 11.8.10. Allow the organic layer to separate from the water phase for a minimum of 10 minutes. If the emulsion interface between layers is more than one-third the volume of the solvent layer, the analyst must employ mechanical techniques to complete the phase separation.
- 11.8.11. Repeat the extraction two additional times with methylene chloride.
- 11.8.12. Determine the original sample volume by re-weighing the sample bottle. Record the sample volume to the nearest centigram (0.01g).
- 11.8.13. Dry extract with sodium sulfate: Place glass wool in a precleaned filter funnel. Rinse glass wool with methylene chloride and load funnel with Na<sub>2</sub>SO<sub>4</sub>. Pour extract through Na<sub>2</sub>SO<sub>4</sub> to remove water. Rinse Na<sub>2</sub>SO with fresh methylene chloride and collect in round bottom flask.
- 11.8.14. Transfer the extract to a 500 mL round-bottom previously labeled with the sample ID, then add approximately 100 μL of tetradecane and concentrate on a rotary evaporator or TurboVap.
- 11.8.15. Perform macro-concentration as detailed in Section 11.17.
- 11.9. Aqueous Samples (solid phase extraction).
  - 11.9.1. Weigh the sample in the bottle on the top loading balance to the nearest centigram (0.01g), and record the mass.
  - 11.9.2. Create a blank and LCS by adding 1L of laboratory reagent water to 2 additional 1L bottles.
  - 11.9.3. For each sample, add 1mL of daily internal standard solution in acetone.

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- Add this solution to the sample in the bottles. Each aliquot of spike mixture is added similarly.
- 11.9.4. To the LCS, add 50µL of the precision and recovery standard in acetone.
- 11.9.5. Prepare the C18 extraction discs by first soaking them in toluene for at least 5 minutes.
- 11.9.6. Assemble the filter holder and vacuum filtration flask and place the extraction disc onto the filter holder. Place a GF-F filter on top of the extraction disc. If the sample has a large amount of particulates a GF-D filter can be placed on top of the GF-F filter. Alternatively, a GMF-150 filter can be used in place of the two filters.
- 11.9.7. Place the filtering funnel onto the disc holder and clamp it in place.
- 11.9.8. Rinse the filter and discs with approximately 15mL of toluene and allow it to soak for about a minute. Apply vacuum and draw the toluene through the discs. Repeat the wash step using about 15mL of acetone. Apply vacuum and draw the acetone through the discs.
- 11.9.9. Rinse the filter and discs with approximately 15mL of methanol and allow it to soak for about a minute. Apply vacuum and draw the methanol through the discs, but **DO NOT ALLOW THE DISCS TO GO DRY**. If they do go dry, simply repeat the methanol rinse step, leaving a 1 2mm layer of solvent on top of the discs.
- 11.9.10. Rinse twice with about 50mL of reagent water, leaving a 1 2mm layer of water on the surface of the discs.
- 11.9.11. Pour the spiked method blank, LCS or sample into the reservoir and apply vacuum to begin the extraction. Adjust the vacuum such that the extraction takes approximately 10 minutes. Samples with large amounts of particulates may take much longer.
- 11.9.12. After most of the sample has been pulled through the discs, rinse the sample bottle with a few mLs of reagent water and add the rinse to the funnel. Rinse down the sides of the funnel with reagent water as well.
- 11.9.13. Allow the discs to dry, remove them from the holder and extract by soxhlet (11.5) or soxtherm (11.6) and proceed with cleanups.
- 11.9.14. Determine the original sample volume by re-weighing the sample bottle. Record the sample volume to the nearest centigram (0.01g).

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# 11.10. Breaking Emulsions

There are several useful methods to decrease or eliminate emulsion in aqueous samples when extracting with methylene chloride. These methods may include stirring with a pipette to manually breakup the emulsions or to transfer the sample into centrifuge tubes and centrifuge at approximately 3000 RPM. The most useful method is to use a 10:1 NaOH/H<sub>2</sub>O solution to change the pH enough to disrupt the emulsion phase, which works 90% of the time. See Section 7.3.5 for reagent preparation.

- 11.10.1. Check the pH of the sample to verify that the pH is between 3 and 7. If the pH is greater than 7, consult the supervisor and client for instructions.
- 11.10.2. Pour approximately 100 mL of the 10:1 NaOH/H<sub>2</sub>O into a 1 L amber glass bottle (AGB).
- 11.10.3. Drain the sample with the emulsion from the 2 L separatory funnel into the 1 L AGB and let it stand.
- 11.10.4. Empty the aqueous waste into the LLE waste drum.
- 11.10.5. Pour the solution with methylene chloride back into the same 2 L separatory funnel and drain the methylene chloride phase through Na<sub>2</sub>SO<sub>4</sub> into a 500 mL round-bottom flask.
- 11.10.6. Empty the aqueous waste into the LLE waste drum.
- 11.10.7. Proceed with macro-concentration (Section 11.17).

# 11.11. Filter/PUF Samples

- 11.11.1. Place the glass sleeve containing the PUF and the Quartz Fiber Filter into the pre-cleaned Soxhlet extractor charged with toluene.
- 11.11.2. Add 2 mL (4000 pg) of 1613/8290 daily Internal Standard solution to all samples and QC.
- 11.11.3. Add 50 uL of 1613/8290 Native Spike to the LCS.
- 11.11.4. Extract the samples and QC for a minimum of 16 hours.
- 11.11.5. Concentrate the extract from the round bottom flask with hexane and adjust the volume.
- 11.11.6. Transfer the extract from the round bottom flask with hexane and adjust the volume.
- 11.11.7. Split the extract 50:50 for analysis and archive.

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## 11.11.8. Proceed to Section 11.12.

# 11.12. Extract Clean-Up

- 11.12.1. For all samples that are not air media, spike 1.0 mL of the Cleanup Recovery Standard (CRS) prior to any cleanup into the round bottom flasks containing the samples and QC Extracts (See also Section 9.7).
- 11.12.2. Proceed with further cleanups as dictated by the sample matrix and extract color. The "Option C" cleanup (Section 11.13) and the IFB Upper Column cleanup (Section 11.14) are applied to samples with high levels of interferences. The IFB column cleanup (Section 11.15) is applied to all samples.

# 11.13. Acid Partitioning ("Option C")

- 11.13.1. Use this clean up as needed on samples with high levels of interferences. Consult with a lead chemist or department manager to determine applicability.
- 11.13.2. Partition the extract in 50-125 mL of hexane against 40 mL concentrated H<sub>2</sub>SO<sub>4</sub> in a separatory funnel. Shake for two minutes. Remove and discard the H<sub>2</sub>SO<sub>4</sub> layer (bottom). Repeat the acid washing until no color is visible in the acid layer (perform a maximum of four acid washings).

Warning: Shaking with a concentrated caustic is a high-risk activity. Analyst must wear a face shield over safety glasses/goggles, or the shaking must take behind a closed hood sash.

11.13.3. Partition the extract against 50 mL of distilled H<sub>2</sub>O. Shake for two minutes. Remove and discard the aqueous layer (bottom). Dry the extract by pouring it through a funnel containing anhydrous sodium sulfate and collect it in a round-bottom flask. Rinse the sodium sulfate with two 15 mL portions of hexane, add the rinsates to the flask, and concentrate the hexane solution to near dryness on a rotary evaporator (35°C water bath), making sure all traces of toluene (when applicable) are removed. (Use of blow-down with an inert gas to concentrate the extract is also permitted.) The DI H<sub>2</sub>O partition is applied only as samples warrant it at the discretion of the analyst.

# 11.14. IFB Upper Column Cleanup

- 11.14.1. Use this clean up as needed on samples with high levels of interferences. Consult with a lead chemist or department manager to determine applicability.
- 11.14.2. Set up the upper of the two chromatography columns as depicted in Figure 2.

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The column (20 mm diameter) is packed in this order: a glass wool plug, 2 g activated silica gel, 4 g Acid silica gel, 2 g activated silica gel, and 1 g sodium sulfate.

- 11.14.3. Pre-rinse the column with 20 mL hexane, and discard the rinsate.
- 11.14.4. Add extract to the column. Rinse extract vessel 2 times with 1 mL each of hexane and add to column.
- 11.14.5. Elute 60 mL hexane directly onto acid silica column (upper column).
- 11.14.6. Collect the eluate, and concentrate before proceeding with the IFB cleanup (Section 11.15).

# 11.15. IFB Column Cleanup

Most samples will undergo this cleanup, either direction following concentration on the rotovap, or following the cleanup in Section 11.13 (Option C) or Section 11.14 (IFB Upper Column).

- 11.15.1. Set up two chromatography columns as depicted in Figure 2. The upper column (20 mm diameter) is packed in this order: a glass wool plug, 2 g activated silica gel, 4 g Acid silica gel, 2 g activated silica gel, and 1 g sodium sulfate. The lower column (15 mm diameter) is packed in this order: a glass wool plug, 6 g acid alumina, and 1 g sodium sulfate.
- 11.15.2. Pre-rinse each column with 20 mL hexane, and discard the rinsate.
- 11.15.3. Put one column above the other.
- 11.15.4. Add extract to the top column (silica column). Rinse extract vessel 2 times with 1 mL each of hexane and add to column.
- 11.15.5. Elute 60 mL hexane directly onto acid silica column (upper column).
- 11.15.6. Discard upper column.
- 11.15.7. Elute lower column with 10 mL of 20% methylene chloride/hexane. Discard in proper waste stream.
- 11.15.8. Elute lower column with 30 mL of 65% methylene chloride/hexane. Save and collect in culture tube.
- 11.15.9. Proceed with additional cleanups as necessary.

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# 11.16. Carbon Column Clean-up (D2 Column)

Prepare an activated Carbon & Silica Gel column as described in below. Refer to the diagram in Figure 3 as well.

- 11.16.1. Push a glasswool plug down to the 3 inch mark in a pre-cut D2 column.
- 11.16.2. Add 1 g of 5% activated carbon/silica. Top with a glasswool plug.
- 11.16.3. With the column oriented with "A" on the top (and the carbon on the lower end of the column), pre-elute with 5 mL 1:1 methylene chloride :cyclohexane.
- 11.16.4. Discard pre-eluates.
- 11.16.5. Invert the column so that the column is oriented with the "B" on the top and pre-elute with 3 mL of 1:1 methylene chloride.
- 11.16.6. Dilute the extract to 1 mL with hexane and transfer to the column (still oriented in the "B" direction).
- 11.16.7. Rinse sample vial onto the column with 2 x 2 mL 1:1 methylene chloride:cyclohexane.
- 11.16.8. Elute with 6 mL 1:1 methylene chloride :cyclohexane
- 11.16.9. Elute with 5 mL 75:25 methylene chloride:methanol
- 11.16.10. Discard eluates.
- 11.16.11. Turn the column over (so that the "A" end is on top), and elute with 30 mL of toluene. Collect this eluate.
- 11.16.12. Concentrate to NEAR dryness using the Rotovap (Section 11.17) or Turbovap (Section 11.18), then proceed to the recovery standard step (Section 11.19).
- 11.17. Macro-concentration (Rotary Evaporator)

Concentrate the extracts in separate round bottom flasks on rotary evaporator.

11.17.1. Assemble the rotary evaporator according to manufacture's instructions, and warm the water bath. On a daily basis, preclean the rotary evaporator by solvent rinsing. Between samples, 2-3 mL rinses of toluene followed by a 2-3 mL rinse of hexane should be rinsed down the feed tube into a waste beaker.

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Rotovap Conditions					
Solvent	Bath Temperature (C)	Vacuum Setting (PSI)			
Toluene	80	25			
Hexane	65	15			
Methylene Chloride	70	No vacuum applied			

- 11.17.2. Attach the round bottom flask containing the sample extract to the rotary evaporator. Slowly apply vacuum to the system, and begin rotating the sample flask.
- 11.17.3. Lower the flask into the water bath and adjust the speed of rotation and the temperature as required. At the proper rate of concentration, the flow of solvent into the receiving flask will be steady, but no bumping or visible boiling of the extract will occur.

*NOTE:* If the rate of concentration is too fast, analyte loss may occur.

- 11.17.4. For samples requiring % Lipids analysis:
  - 11.17.4.1. Concentrate until the toluene has been completely removed. Add approximately 25 mL hexane and concentrate to ensure that only the lipids are present.
  - 11.17.4.2. Dry the concentration vessel and let stand at room temperature. Weigh the vessel and record on the benchsheet.
  - 11.17.4.3. Calculate % lipids as follows:

$$\% \ Lipids = \frac{Final \ Vessel \ Mass - Initial \ Vessel \ Mass}{Sample \ Size} \times 100\%$$

- 11.17.5. Proceed to extract cleanups, or transfer to a micro concentration vial for the recovery standard step (Section 11.19).
- 11.18. Micro-concentration (Turbovap)

Concentrate the extracts in 35 mL culture tubes in a turbo-evaporator. The turbo-evaporator model that the laboratory uses can hold up to 50-35 mL culture tubes. Other turbo-evaporator models can be used that may or may not have the same culture tube sizes and/or capacity. Adjust temperature according to solvent (65°C for toluene and 45°C for hexane or hexane/ methylene chloride mixtures)

- 11.18.1. The evaporating times are dependent on sample volume and solvent. The following are examples and can change from sample to sample. Each sample should be checked in intermittent intervals to make sure samples do not go dry.
- 11.18.2. When evaporating 30 mL toluene, it will normally take approximately 30-50

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minutes with the temperature setting described above.

- 11.18.3. When evaporating 30 mL hexane/ methylene chloride, it will normally take approximately 20-30 minutes with the temperature setting described above.
- 11.18.4. For samples requiring % Lipids analysis refer to Section 11.17.4.
- 11.18.5. Proceed to extract cleanups, or transfer to a micro concentration vial for the recovery standard step (Section 11.19).

# 11.19. Recovery Standard

- 11.19.1. Transfer extracts to a micro concentration vial (test tubes and other small vessels may also be used)
- 11.19.2. With a stream of dry, purified nitrogen, reduce the extract volume to approximately  $100 \mu L$ .
- 11.19.3. Add 20 µL of the recovery standard solution (Table 2).
- 11.19.4. With a stream of dry, purified nitrogen, reduce the extract volume to 20 μL.
- 11.19.5. Transfer the extract to an autoinjection vial and store in the dark at room temperature.
- 11.19.6. A smaller final volume can be used to decrease the detection limit upon client approval.
- 11.19.7. A larger final volume can be use to decrease potential matrix interferences, if the column and acid cleanups were unsuccessful.

# 11.20. Sample Dilution Procedure

11.20.1. Simple dilutions: Dilutions from 2X to 50X can be achieved without respiking the final extract. The calculation to determine the final extract concentration is as follows:

Final Conc. of Extract = 
$$\frac{\text{(Conc. of original extract)} \times \text{(Amount of aliquot taken)}}{\text{(Volume of diluted extract)}}$$

Ex: 
$$\frac{(10 \text{ g}) \text{ x} (2 \mu \text{L})}{(20 \mu \text{L}) \text{ x} (100 \mu \text{L})} = \frac{1 \text{ g}}{100 \mu \text{L}} \text{ FV}$$

Record the final sample concentration on the extract label.

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# 11.20.2. Complex dilution requiring respiking of IS and RS:

Dilutions greater than 50x must be done by diluting and respiking the extract with IS and RS. This procedure may require serial dilution to be performed. If this procedure is done, then the sample size must be adjusted to reflect the aliquot taken.

Ex. 100X dilution (original sample with 10 g/20 μL final volume)

Take a 2  $\mu$ L aliquot (1/10 of original sample) and add 18  $\mu$ L of solvent keeper. Take a 2  $\mu$ L aliquot of the dilution (1/100 of the original sample), respike with 1 mL IS and 20  $\mu$ L RS, reduced to 20  $\mu$ L FV.

Record the final sample concentration of the extract label.

## 12. CALCULATIONS/DATA REDUCTION

12.1. Not applicable

## 13. METHOD PERFORMANCE

It must be documented that all applicable system performance criteria specified were met before analysis of any sample is performed.

13.1. The group/team leader has the responsibility to ensure that this procedure is performed by an associate who has been properly trained in its use and has the required expertise.

#### 13.2. Method Detection Limit

The laboratory must generate a valid method detection limit for each analyte of interest. The MDL must be below the reporting limit for each analyte. The procedure for determination of the method detection limit is given in 40 CFR Part 136, Appendix B, and further defined in SOP WS-QA-0006. MDLs are available in the Quality Assurance Department.

#### 13.3. Initial Demonstration

The laboratory must make an initial demonstration of capability for each individual method. Demonstration of capability for both soil and water matrices is required. This requires analysis of QC check samples containing all of the standard analytes for the method. For some tests it may be necessary to use more than one QC check mix to cover all analytes of interest.

- 13.3.1. Four aliquots of the QC check sample are analyzed using the same procedures used to analyze samples, including sample preparation. The concentration of the QC check sample should be less than or equivalent to the LCS samples.
- 13.3.2. Calculate the average recovery and standard deviation of the recovery for each analyte of interest. Compare these to the laboratory generated QC Limits.

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13.4. If any analyte does not meet the acceptance criteria the test must be repeated. Only those analytes that did not meet criteria in the first test need to be evaluated. Repeated failure for any analyte indicates the need for the laboratory to evaluate the analytical procedure and take corrective action.

## 14. POLLUTION CONTROL

It is TestAmerica's policy to evaluate each method and look for opportunities to minimize waste generated (i.e., examine recycling options, ordering chemicals based on quantity needed, preparation of reagents based on anticipated usage and reagent stability). Employees must abide by the policies in Section 13 of the Corporate Environmental Health and Safety Manual (CW-E-M-001) for "Waste Management and Pollution Prevention."

- 14.1. The use of Roto-vaps and Turbo-vaps rather than Kuderna-Danish reduction allows extraction solvents to be collected and disposed of rather than released to the atmosphere.
- 14.2. Toluene, which is a less hazardous solvent, has been substituted for benzene as an extraction solvent.
- 14.3. The use of SoxTherm extraction rather than soxhlet extraction, when appropriate, reduces the volume of solvent used.
- 14.4. Standards should be prepared in volumes consistent with laboratory use to minimize the volume of expired standards that must be discarded.
- 14.5. All waste will be disposed of in accordance with Federal, State, and Local regulations. Where reasonably feasible, technological changes have been implemented to minimize the potential for pollution of the environment.
- 14.6. Do not allow waste solvent to vent into the hoods. All solvent waste is stored in capped containers unless they are being filled.
- 14.7. Transfer waste solvent from collection cups (tri-pour and similar containers) to jugs and/or carboys as quickly as possible to minimize evaporation.

## 15. WASTE MANAGEMENT

Waste management practices are conducted consistent with all applicable rules and regulations. Excess reagents, samples and method process wastes are disposed of in an accepted manner. Waste description rules and land disposal restrictions are followed. Waste disposal procedures are incorporated by reference to SOP WS-EHS-0001. The following waste streams are produced when this method is carried out.

15.1. Extracted aqueous/leachate samples contaminated with methylene chloride are collected at the fume hood in a 5-gallon or smaller carboy. If the samples are not at a

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neutral pH, add small quantities of sodium bicarbonate to bring the waste to neutral. Stir well. Once neutralized, immediately pour the carboy contents into a blue plastic LLE drum in the H3 closet. When full to between two and six inches of the top, or after no more than 75 days, move the LLE drum to the waste collection area for shipment.

- 15.2. Extracted soil samples and thimbles, extracted PUF filters, XAD-2 resin, paper funnel filters, glass wool, sodium sulfate, assorted disposable glassware, fish/crawfish or similar materials, silica gel, alumina, and carbon from column clean-ups, contaminated with various solvents and eluates. Dump the materials into a orange contaminated lab trash bucket. When the bucket is full or at the end of the day, tie the plastic bag liner shut and put the lab trash into the steel collection drum in the H3 closet. When the drum is full or after no more than 75 days, move it to the waste collection area for shipment.
- 15.3. Flammable solvent and methylene chloride waste generated during glassware and sodium sulfate cleaning. Solvent waste collected during roto-vap/turbo-vap reduction of extracted samples. Collect the waste solvents in tripours during use. Empty the tripours into a 1-liter to 4-liter carboy at the fume hood. When the carboy is full, or at the end of your shift, whichever comes first, empty the carboy into the steel solvent drum in the H3 closet. When full to between two and six inches of the top, or after no more than 75 days, move the steel drum to the waste collection area for shipment.
- 15.4. Assorted flammable solvents and methylene chloride waste generated during quartz fiber filter preparation, PUF adsorbent preparation, XAD-2 resin preparation, PUF/XAD-2 cartridge preparation, glassware rinsing and sodium sulfate pre-rinsing. Waste solvents and methylene chloride collected during roto-rap/turbo-vap reduction of extracted samples. Collect the waste solvents in tripours during use. Empty the tripours into a 1-liter to 4-liter carboy at the fume hood. When the carboy is full, or at the end of your shift, whichever comes first, empty the carboy into the steel drum in the H3 closet. When the drum is full to between two and six inches of the top, or after no more than 75 days, move the steel drum to the waste collection area for shipment.
- 15.5. Contaminated sulfuric acid used during extract cleanup. Collect the used sulfuric acid in empty, 2.5-liter, plastic coated jars. When full or after one year, whichever comes first, transfer these jars to the waste collection area for shipment.
- 15.6. Contaminated distilled water used during extract cleanup. Collect the contaminated water in a 1-liter to 4-liter carboy at the fume hood. When the carboy is full, or at the end of your shift, whichever comes first, empty the carboy into the plastic LLE drum in the H3 closet. When full to between two and six inches of the top, or after no more than 75 days, move the plastic drum to the waste collection area for shipment.

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## 16. REFERENCES/CROSS REFERENCES

- 16.1. SW846, Test Methods for Evaluating Solid Waste, Third edition, Update IV. Method 8290A Polychlorinated Dibenzodioxins (PCDDs) and Polychlorinated Dibenzofurans (PCDFs) by high-Resolution Mass Spectrometry February 2007.
- 16.2. SW846, Test Methods for Evaluating Solid Waste, Third edition, Update III. Method 8290 Polychlorinated Dibenzodioxins (PCDDs) and Polychlorinated Dibenzofurans (PCDFs) by high-Resolution Mass Spectrometry September 1994.
- 16.3. SW846, Test Methods for Evaluating Solid Waste, Third edition, Update III. Method 0023A, Sampling Method for Polychlorinated Dibenzo-p-dioxins and Polychlorinated Dibenzofurans Emissions from Stationary Sources. December 1996.
- 16.4. Compendium Method TO-9A "Determination of Polychlorinated, Polybrominated, and Brominated, Cholorinated Dibenxo-p-dioxins and Dibenzofurans in Ambient Air", EPA compendium of Methods for the Determination of Toxic Organic Compounds in Ambient Air, second edition, January 1997.
- 16.5. Protocol for the Analysis of 2,3,7,8-TCDD by HRGC/HRMS". J. S. Stanley and T. M. Sack, EPA 600/4-86-004.
- 16.6. "Safety in Academic Chemistry Laboratories", American Chemical Society Publication, Committee on Chemical Safety (3rd Edition, 1979.)
- 16.7. "Carcinogens Working with Carcinogens". Department of Health, Education, and Welfare, Public Health Service, Center for Disease Control. National Institute for Occupational Safety and Health. Publication No. 77-206, August 1977.
- 16.8. "OSHA Safety and Health Standards, General Industry", (29 CFR 1910) Occupational Safety and Health Administration, OSHA 2206 (revised January 1976).

### 17. METHOD MODIFICATIONS

- 17.1. Deviations from EPA 8290 and 8290A.
  - 17.1.1. Tetradecane instead of nonane is used as the final solvent to increase the stability of extracts and standards. Tetradecane is less volatile than nonane. Loss of analyte as a result of solvent incompatibility is monitored through recovery checks and calibration acceptance criteria.
  - 17.1.2. Extract clean-ups are performed at the discretion of the analyst when interferences are observed. Then, the analyst should select the clean-up procedure appropriate to the interferent.

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- 17.1.3. Section 7.4.6.4 of Method 8290 indicates that extracts should be transferred with hexane, then toluene. Toluene is used to transfer extracts to maintain compound solubility and minimize analyte loss.
- 17.1.4. Section 7.5.1.2 of Method 8290 specifies that a NaCl solution should be used for partitioning. Instead, the laboratory uses laboratory water only. NaCl is used to break up emulsions that may form. An analyst may use NaCl, NaOH, or any mechanical means to break up an emulsion.
- 17.1.5. Section 7.5.3 of Method 8290 specifies that hexane is used as a column elution solvent. The laboratory uses cyclohexane to achieve better and more reproducible separation of the target analyte from the interferent.
- 17.1.6. Carbon columns are packed with silica gel in place of celite. Elution solvents are changed accordingly. (SOP Section 11.4; Method 8290 Section 7.5.3.2, 8290A Section 7.3.6.).

## 17.2. Modifications from TO-9A method

- 17.2.1. Quartz Fiber Filters are cleaned by Soxhlet extraction with methylene chloride, not baked at 400 degrees C for 5 hours.
- 17.2.2. The PUF material may be pre-cleaned with methylene chloride or other appropriate solvent. The PUFs are not reused.
- 17.2.3. The  $^{37}\text{Cl}_4$ -2,3,7,8-TCDD surrogate is present at varying levels in the calibration curve (0.5-200 pg/  $\mu$ L).
- 17.2.4. Samples are extracted with toluene not benzene.
- 17.2.5. Concentration is performed by rotary evaporation not Kuderna-Danish.
- 17.2.6. All cleanup procedures are optional and applied based on the analyst's discretion.
- 17.2.7. The laboratory uses 2 labeled recovery standard for the quantitation of labeled internal standards.
- 17.2.8. The final volume is adjusted to 20 µL in tetradecane.
- 17.2.9. Calibration and quantitation are performed in accordance to this SOP.

# 18. ATTACHMENTS

18.1. Table 1 - Types of Matrices

- 18.2. Table 2 Composition of Sample Fortification and Recovery Standard Solutions.
- 18.3. Table 3 The Seventeen 2,3,7,8-Substituted PCDD and PCDF Congeners
- 18.4. Figure 1 Analysis Flowchart
- 18.5. Figure 2 IFB column cleanup
- 18.6. Figure 3 D2 Column cleanup
- 18.7. Appendix A Periodic Wipe Test Performance

## 19. REVISION HISTORY

- 19.1. WS-IDP-0005, Revision 1.5, Effective 12/21/2012
  - 19.1.1. Clarified extraction procedure by revising Section(s) 11.8.1- 11.8.4 and adding an extra extraction step (Section 11.8.3).
  - 19.1.2. Editorial revisions.
- 19.2. WS-IDP-0005, Revision 1.4, Effective 03/20/2012
  - 19.2.1. Appended to Section 2.2: "This method can also use solid phase extraction (SPE), however, Test America West Sacramento is in the developmental stages for this extraction type and is not currently certified for its use."
  - 19.2.2. Editorial changes.
- 19.3. WS-IDP-0005, Revision 1.3., Effective 06/10/2011
  - 19.3.1. Added Section 11.9: Aqueous Samples (Solid Phase Extraction).
  - 19 3 2 Editorial revisions
- 19.4. WS-IDP-0005, Revision 1.2, Effective 2/11/2011
  - 19.4.1. Added benzene to Section 5.2 Table...
  - 19.4.2. Editorial revisions.
- 19.5. WS-IDP-0005, Revision 1.1, Effective 2/12/2010
  - 19.5.1. Section 11.2 updated SOP reference from SAC-ID-0009 to WS-ID-0009.
  - 19.5.2. Section 11.6.1 changed: "Prior to loading samples, run the system through

- a cleaning cycle (approximately 3 hours)" to "(approximately 1 hour)."
- 19.5.3. Section 11.6.8 changed "...fresh charge (140 mL) of toluene..." to "...fresh charge (150 mL) of toluene...".
- 19.5.4. Section 11.16.1 inserted in Table "No vacuum applied" under vacuum setting (PSI) for solvent Methylene chloride.
- 19.6. WS-IDP-0005, Revision 1, Effective 10/2/2008
  - 19.6.1. Added 8290A references.
    - 19.6.1.1. Extract and standard storage.
    - 19.6.1.2. Removal of MS/MSD.
  - 19.6.2. Updated to TestAmerica format.
  - 19.6.3. Separated the analytical steps from the preparation steps, this SOP is concerned only with the sample preparation.
- 19.7. WS-ID-0005, Revision 6.7, Effective 8/21/2008
  - 19.7.1. Changed the word "toluene" to "acetone" in 7.11.2.
- 19.8. WS-ID-0005, Revision 6.6, Effective 4/9/2008
  - 19.8.1. Added South Carolina rule to prepare an MS/MSD with every batch.
  - 19.8.2. Modified to include extraction and analysis of ambient air samples collected in filter/PUF material.

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TABLE 1

Types of Matrices, Sample Sizes and 2,3,7,8-TCDD-Based
Method Calibration Limits (Parts per Trillion)

	Water	Soil Sediment Paper Pulp	Fly Ash	Human/ Fish Tissue	Adipose Tissue	Sludges, Fuel Oil	Still- Bottom	Ambient or Source Samples
Lower MCL(a)	0.01	1.0	2.0	1.0	2.0	10	20	40
Upper MCL(a)	4.0	400	400	400	400	2000	4000	8000
Weight (g)	1000	10	10	10	10	2.0	1.0	1 sample
IS Spiking Levels (ng)	2.0	2.0	2.0	2.0	2.0	2.0	2.0	4.0
Final Extract Volume (μL)	20	20	20	20	20	20	20	20

<sup>(</sup>a) For other congeners, multiply the values by 1 for TCDF, by 5 for PeCDD/PeCDF/HxCDD/HxCDF/HpCDD/HpCDF, and by 10 for OCDD/OCDF.

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TABLE 2

Composition of the Sample Fortification and Recovery Standard Solutions

Analyte	Semple Fortification Solution	Recovery Standard Solution Concentration pg/µL; Solvent:		
-	Concentration pg/μL;			
	Solvent: Isooctane	Tetradecane		
<sup>13</sup> C <sub>12</sub> -2,3,7,8-TCDD	2 <sup>(a)</sup> , 100 <sup>(c)</sup>			
<sup>13</sup> C <sub>12</sub> -2,3,7,8-TCDF	2 <sup>(a)</sup> , 100 <sup>(c)</sup>			
<sup>13</sup> C <sub>12</sub> -1,2,3,4-TCDD		100		
<sup>13</sup> C <sub>12</sub> -1,2,3,7,8-PeCDD	2 <sup>(a)</sup> , 100 <sup>(c)</sup>			
<sup>113</sup> C <sub>12</sub> -1,2,3,7,8-PeCDF	2 <sup>(a)</sup> , 100 <sup>(c)</sup>			
<sup>13</sup> C <sub>12</sub> -1,2,3,6,7,8-HxCDD	2 <sup>(a)</sup> , 100 <sup>(c)</sup>			
<sup>13</sup> C <sub>12</sub> -1,2,3,4,7,8-HxCDF <sup>(d)</sup>	2 <sup>(a)</sup> , 100 <sup>(c)</sup>			
<sup>113</sup> C <sub>12</sub> -1,2,3,7,8,9-HxCDD		100		
<sup>13</sup> C <sub>12</sub> -2,3,7,8-TCDD <sup>(b)(c)</sup>	0.8 <sup>(b),</sup> 100 <sup>(c)</sup>			
	100 <sup>(c)</sup>			
<sup>13</sup> C <sub>12</sub> -2,3,4,7,8-PeCDF <sup>(c)</sup>	100 <sup>(c)</sup>			
<sup>13</sup> C <sub>12</sub> -1,2,3,6,7,8-HxCDF <sup>(c)(d)</sup>	100 <sup>(c)</sup>			
<sup>13</sup> C <sub>12</sub> -1,2,3,4,7,8-HxCDD <sup>(c)</sup>	100 <sup>(c)</sup>			
<sup>13</sup> C <sub>12</sub> -1,2,3,4,7,8,9-HpCDD <sup>(c)</sup>	100 <sup>(c)</sup>			
<sup>13</sup> C <sub>12</sub> -1,2,3,4,6,7,8-HpCDD	2 <sup>(a)</sup> , 100 <sup>(c)</sup>			
<sup>13</sup> C <sub>12</sub> -1,2,3,4,6,7,8-HpCDF	2 <sup>(a)</sup> , 100 <sup>(c)</sup>			
<sup>13</sup> C <sub>12</sub> -OCDD	4 <sup>(a)</sup> , 200 <sup>(c)</sup>			
3 <sub>12</sub> 3355	7 , 200			

- (a) Standard 8290, Method 23, Method 0023A, TO9 and TO9A Sample Fortification Solution concentrations
- (b) Method TO9 and TO9A surrogate concentrations
- (c) Method 23 and Method 0023A surrogate concentrations
- (d)  $^{13}C_{12}$  -1,2,3,6,7,8-HxCDF is used as a Sample Fortification Solution and  $^{13}C_{12}$ -1,2,3,4,7,8-HxCDF is used as a surrogate solution in Method 23 and Method 0023A

**TABLE 3** The Seventeen 2,3,7,8-Substituted PCDD and PCDF Congeners

PCDD	PCDF
2,3,7,8-TCDD(*)	2,3,7,8-TCDF(*)
1,2,3,7,8-PeCDD(*)	1,2,3,7,8-PeCDD(*)
1,2,3,6,7,8-HxCDD(*)	2,3,4,7,8-PeCDF
1,2,3,4,7,8-HxCDD	1,2,3,6,7,8-HxCDF
1,2,3,7,8,9-HxCDD(+)	1,2,3,7,8,9-HxCDF
1,2,3,4,6,7,8-HpCDD(*)	1,2,3,4,7,8-HxCDF(*)
1,2,3,4,5,6,7,8-OCDD(*)	2,3,4,6,7,8-HxCDF
	1,2,3,4,6,7,8-HpCDF(*)
	1,2,3,4,7,8,9-HpCDF
	1,2,3,4,5,6,7,8-OCDF

<sup>(\*)</sup>The <sup>13</sup>C -labeled analog is used as an internal standard. (+)The <sup>13</sup>C -labeled analog is used as a recovery standard.

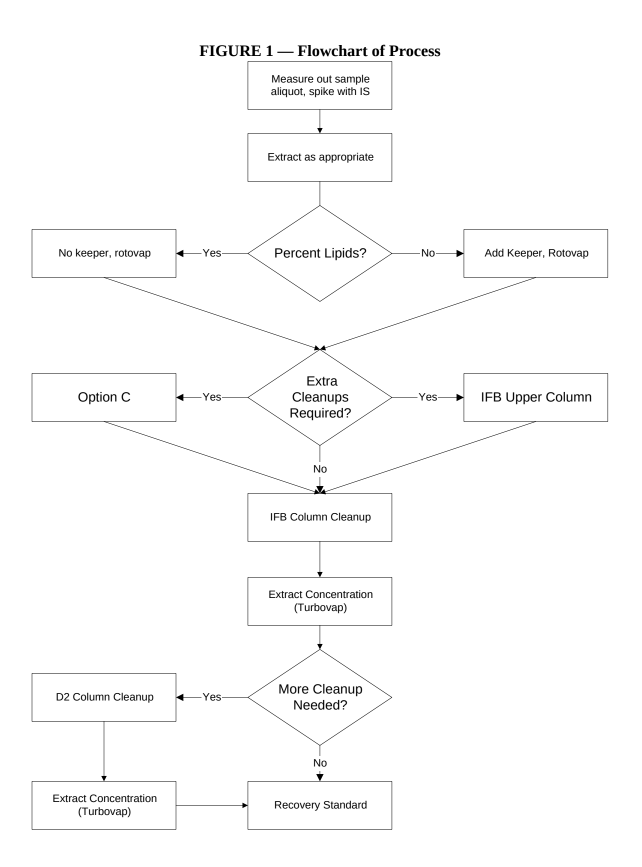


Figure 2 – Diagram of IFB Column Cleanup

Use 20 mm column for top column (IFB Column)

Use 16 mm column for bottom column\* (Acid Alumina)

Note: Upper and lower columns are piggy backed for IFB cleanup, upper column only can be used for additional cleaning.

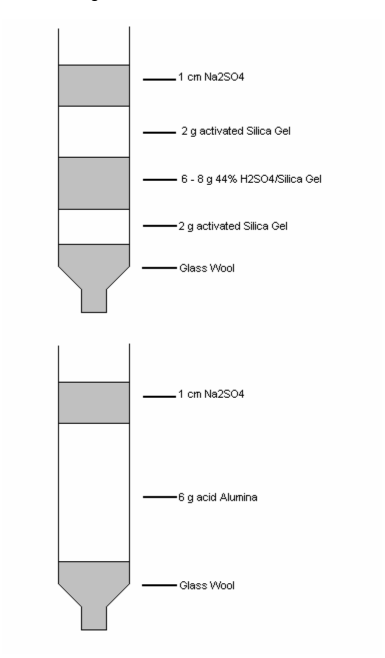
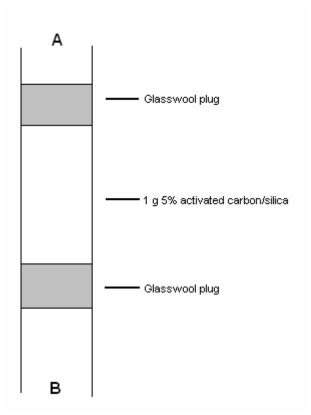


Figure 3— D2 Carbon Column:



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# **APPENDIX A** — Screening the Laboratory for 2,3,7,8 Congeners

This procedure is designed for the periodic evaluation of potential contamination by 2,3,7,8-substituted PCDD/PCDF congeners of the working areas inside the laboratory.

## PERFORMING WIPE TEST

Perform the wipe tests on surface areas of two inches by one foot with laboratory wipers saturated with distilled-in-glass acetone or appropriate solvent using a pair of clean stainless steel forceps. Use one wiper for each of the designated areas. Combine the wipers to one composite sample in an extraction jar containing 200 mL distilled-in-glass hexane. Place an equal number of unused wipers in 200 mL hexane and use this as a control.

## SAMPLE PREPARATION

Close the jar containing the wipes and 200 mL hexane and extract for 20 minutes using a wrist-action shaker. Use an appropriate means to reduce the volume to approximately 1.0 mL. Put through an alumina column to clean up potential interfering compounds. Add appropriate amount of recovery standard.

# **EXTRACT ANALYSIS**

Concentrate the contents of the vial to a final volume of  $20 \mu L$  (either in a minivial or in a capillary tube). Inject  $2 \mu L$  of each extract (wipe and control) onto a capillary column and analyze for 2,3,7,8-substituted PCDDs/PCDFs as specified in the analytical method Section 11 (this exhibit). Perform calculations according to Section 12 (this exhibit).

## REPORTING FORMAT

Report the presence of 2,3,7,8-substituted PCDDs and PCDFs as a quantity (pg or ng) per wipe test experiment (WTE). Under the conditions outlined in this analytical protocol, a lower limit of calibration of 25 pg/WTE is expected for 2,3,7,8-TCDD. A positive response for the blank (control) is defined as a signal in the TCDD retention time window at any of the masses monitored which is equivalent to or above 8 pg of 2,3,7,8-TCDD per WTE. For other congeners, use the multiplication factors listed in Table 1, footnote (a) (e.g., for OCDD, the lower MCL is  $25 \times 5 = 125 \text{ pg/WTE}$  and the positive response for the blank would be  $8 \times 5 = 40 \text{ pg}$ ). Also, report the recoveries of the internal standards during the simplified cleanup procedure.

## FREQUENCY OF WIPE TESTS

Wipe tests should be performed when there is evidence of contamination in the method blanks.

# CORRECTIVE ACTION

An upper limit of 25 pg per TCDD isomer and per wipe test experiment is allowed. (Use multiplication factors listed in footnote (a) from Table 1 for other congeners.) This value corresponds to the lower calibration limit of the analytical method. Steps to correct the contamination must be taken whenever these levels are exceeded. To that effect, first vacuum the working places (hoods, benches, sink) using a vacuum cleaner equipped with a high-efficiency

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particulate absorbent (HEPA) filter and then wash with a detergent. A new set of wipes should be analyzed before anyone is allowed to work in the dioxin area of the laboratory.

The test results and the decontamination procedure must be reviewed with EH&S.